## EST AVAILABLE COT

## (12) DEMANDE INTERNATIONALE PUBLIÉE EN VERTU DU TRAITÉ DE COOPÉRATION EN MATIÈRE DE BREVETS (PCT)

(19) Organisation Mondiale de la Propriété Intellectuelle Bureau international





(43) Date de la publication internationale 21 novembre 2002 (21.11.2002)

**PCT** 

(10) Numéro de publication internationale WO 02/092818 A2

- (51) Classification internationale des brevets<sup>7</sup>:

  C12N 15/31, 15/62, 15/74, 5/10,
  1/20, 1/21, C07K 14/315, 16/12, 19/00, C12Q 1/68, G01N
  33/53, A61K 48/00, 39/395, 39/09, A61P 31/04
- (21) Numéro de la demande internationale :

PCT/IB02/03059

- (22) Date de dépôt international : 26 avril 2002 (26.04.2002)
- (25) Langue de dépôt :

français

(26) Langue de publication :

français

- (30) Données relatives à la priorité : 2(01)/05642 26 avril 2001 (26.04.2001) F
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- (81) États désignés (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW.
- (84) États désignés (régional): brevet ARIPO (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), brevet eurasien (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), brevet européen (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), brevet OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

## Publiée:

- sans rapport de recherche internationale, sera republiée des réception de ce rapport
- avec la partie réservée au listage des séquences de la description publiée séparément sous forme électronique et disponible sur demande auprès du Bureau international

En ce qui concerne les codes à deux lettres et autres abréviations, se référer aux "Notes explicatives relatives aux codes et abréviations" figurant au début de chaque numéro ordinaire de la Gazette du PCT.

(54) Title: STREPTOCOCCUS AGALACTIAE GENOME SEQUENCE, USE FOR DEVELOPING VACCINES, DIAGNOSTIC TOXILS. AND FOR IDENTIFYING THERAPEUTIC TARGETS

(54) Titre: SEQUENCE DU GENOME STREPTOCOCCUS AGALACTIAE, APPLICATION AU DEVELOPPEMENT DE VAC-CINS, D'OUTILS DE DIAGNOSTIC, ET A L'IDENTIFICATION DE CIBLES THERAPEUTIQUES

(57) Abstract: The invention concerns the genome sequence and nucleotide sequences coding for Streptococcus agalactiae polypeptides, such as cellular envelope polypeptides, or secreted or specific polypeptides, or polypeptides involved in the metabolism and the replication process, as well as vectors or cells comprising said sequences. The invention also concerns the use thereof for developing vaccines, diagnostic tools, DNA chips and for identifying therapeutic targets.

(57) Abrégé: L'invention a pour objet la séquence génomique et des séquences nucléotidiques codant pour des polypeptides de Streptococcus agalactiae, tels que des polypeptides d'enveloppe cellulaire, ou des polypeptides sécrétés ou spécifiques, ou impliqués dans le métabolisme et dans le processus de réplication, ainsi que des vecteurs ou cellules incluant lesdites séquences. L'invention porte également sur leur application au développement de vaccins, d'outils de diagnostic, de puces à ADN et à l'identification de cibles thérapeutiques.

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Séquence du génome *Streptococcus agalactiae*, application au développement de vaccins, d'outils de diagnostic, et à l'identification de cibles thérapeutiques.

L'invention a pour objet la séquence génomique et des séquences nucléotidiques codant pour des polypeptides de *Streptococcus agalactiae*, tels que des polypeptides d'enveloppe cellulaire, ou des polypeptides sécrétés ou spécifiques, ou impliqués dans le métabolisme et dans le processus de réplication, ainsi que des vecteurs ou cellules incluant lesdites séquences. L'invention porte également sur leur application au développement de vaccins, d'outils de diagnostic, de puces à ADN et à l'identification de cibles thérapeutiques.

Streptococcus agalactiae est un streptocoque \( \beta\)-hémolytique qui constitue la seule espèce appartenant au groupe B de Lancefield (SGB). Les infections néonatales à streptocoques du groupe B posent un important problème de santé publique qui ne se limite pas aux pays en voie de développement. Leur incidence est de 2,5 pour 1 000 naissances, avec un taux de mortalité qui varie actuellement dans les pays industrialisés entre 4 et 10 % selon les études. Cette bactérie est responsable d'environ 20 % des méningites bactériennes recensées en France et des séquelles neurologiques sont alors observées dans 25 à 50 % des cas. Elle est également à l'origine de mort foetale in utero. Le polyoside capsulaire est l'antigène de surface majeur des SGB. Cinq sérotypes (Ia, lb, II, III et V) sont généralement détectés au cours des infections humaines, le sérotype III étant retrouvé dans 75 % des infections néonatales avec atteinte méningée. Au niveau cellulaire et moléculaire, les différentes étapes du processus infectieux dû à S. agalactiae sont encore peu connues. Il est vraisemblable que, dans le cas du syndrome précoce (infections survenant dans les 24 premières heures), la bactérie inhalée pénètre dans les cellules de l'épithélium alvéolaire du nouveau-né et traverse cette barrière pour disséminer ultérieurement dans la circulation générale. La genèse du syndrome tardif précoce (infections survenant entre le 7ème jour et le 3ème mois) et des autres infections à SGB reste encore très mal comprise. Le seul facteur de virulence des SGB dont le rôle a été clairement démontré est le polyoside capsulaire qui permet l'échappement au système immunitaire de l'hôte. La contribution exacte de certaines protéines de surface (antigène C, protéine Rib et C5a peptidase) à la virulence de cette bactérie est encore peu connue.

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Une recherche réalisée sur le site EXPASY (http://www.expasy.ch/) indique qu'il existe 112 références de séquences protéiques dans les banques Swissprot et TREMBL. Ce nombre inclut des protéines codées par des plasmides de S. agalactiae. Ces séquences représentent donc une vision partielle d'un nombre limité d'aspect de la biologie de S. agalactiae. La biosynthèse de la capsule polysaccharidique est un des aspects les mieux connus de la virulence de cette bactérie. Par ailleurs, les gènes codant pour 6 protéines exposées à la surface sont également connus (3).

Afin d'appréhender de manière globale les déterminants génétiques impliqués dans ces processus ainsi que le métabolisme de Streptococcus agalactiae, le séquençage du génome de Streptococcus agalactiae a été réalisé. Le génome de la souche Streptococcus agalactiae CIP 82.45 (ATCC 12403) qui a été responsable d'une septicémie mortelle a été choisi pour ce séquençage. Cette souche possède un sérotype capsulaire III, ne présente pas de résistance acquise aux antibiotiques, est génétiquement modifiable et est virulente dans un modèle d'infection murin. La connaissance complète du génome est une étape cruciale pour la caractérisation des gènes impliqués dans le développement du processus infectieux : adhésion et franchissement des structures épithéliales, échappement au système immunitaire et adaptation à des conditions de culture variées et souvent hostiles (pH, stress oxydatif et carences nutritionnelles), qui constituent des cibles potentielles pour de nouvelles stratégies thérapeutiques. La comparaison du génome de S. agalactiae avec ceux d'autres pathogènes à Gram positif (Streptococcus pyogenes, Streptococcus pneumoniae, Streptococcus Staphylococcus aureus, Listeria monocytogenes, ...) doit permettre d'identifier de nouveaux gènes de virulence ainsi crue nouvelles cibles pour construire des souches de virulence atténuées et des vaccins. Les protéines de surfaces constituent des candidats pour une future préparation vaccinale. Dans les tableaux 2 et 6 ci-après sont répertoriés respectivement 25 et 30 nouveaux gènes, nouvellement identifiés, codant pour des protéines potentiellement liées au peptidoglycane et présentant le motif de liaison LPXTG.

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La séquence complète du génome de *Streptococcus agalactiae* (CIP 82.45 (ATCC 12403)) a été obtenue. Ce génome est constitué d'un chromosome long d'environ 2,2 Mb identifié ici sous forme de 138 contigs représentés par les séquences SEQ ID No. 1 à SEQ ID No. 136, SEQ ID No. 138 et SEQ ID No. 139, et d'un plasmide

long de 45 kbases présent dans la souche séquencée représenté par la séquence SEQ ID No. 137. La séquence complète du génome est représentée par la séquence SEQ ID No. 2345.

Une liste des phases codantes annotées identifiées par l'analyse des séquences de ces contigs est donnée au tableau 1.

Une liste des phases codantes pour des protéines de surface nouvellement identifiées est donnée au tableau 2 comme indiqué précédemment.

Une liste des phases codantes annotées identifiées par l'analyse de la séquence génomique complète SEQ ID No. 2345 est donnée au tableau 3.

Une liste des phases codantes pour des protéines de surface identifiées à partir de l'analyse de la séquence génomique complète SEQ ID No. 2345 est donnée au tableau 6 (protéines liées au peptidoglycane), tableau 8 (lipoprotéines). Tableau 9 (autres protéines de suface), Tableau 10( protéines impliquées dans la biosynthèse des composés polysaccharidiques)

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La présente invention concerne les séquences nucléotidiques et polypeptidiques de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403).

Ainsi, c'est un objet de la présente invention que de caractériser la séquence du génome de *Streptococcus agalactiae*, CIP 82.45 (ATCC 12403) contenu dans la banque génomique préparée à partir du génome de cette souche et déposée à la CNCM le 28 décembre 2000 sous le numéro I-2610, ainsi que de tous les gènes et séquences régulatrices non codantes contenus dans ledit génome.

La présente invention concerne donc une séquence nucléotidique isolée et/ou purifiée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi les séquences SEQ ID No. 1 à SEQ ID No. 139 et la séquence SEQ ID No. 2345.

La présente invention concerne également une séquence nucléotidique isolée et/ou purifiée, issue de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi :

- a) une séquence nucléotidique comportant au moins 75 %, 80 %, 85 %, 90 %, 95 % ou 98 % d'identité avec une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345;
- b) une séquence nucléotidique hybridant dans des conditions de forte stringence avec une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345,

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et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides;

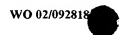
- c) une séquence nucléotidique complémentaire d'une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, ou complémentaire d'une séquence nucléotidique telle que définie en a), ou b), ou une séquence nucléotidique de l'ARN correspondant à l'une des séquences a) ou b);
- d) une séquence nucléotidique d'un fragment représentatif d'une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, ou d'un fragment représentatif d'une séquence nucléotidique telle que définie en a), b) ou c) et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides;
- e) une séquence nucléotidique comprenant une séquence telle que définie en a), b), c) ou d); et
- f) une séquence nucléotidique telle que définie en a), b), c), d) ou e) modifiée, de préférence comportant au plus 10 %, 5 %, 1 % ou 0, 5 % de nucléotides modifiés par rapport à la séquence de référence.

De façon plus particulière, la présente invention a également pour objet les séquences nucléotidiques isolées et/ou purifiées, caractérisées en ce qu'elles sont issues de SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, et en ce qu'elles codent pour un polypeptide choisi parmi les polypeptides de séquence SEQ ID No. 140 à SEQ ID No. 2344, et SEQ ID No. 2346 à SEQ ID No. 4481.

La présente invention concerne aussi de façon plus générale les séquences nucléotidiques issues de SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, et codant pour un polypeptide de *Streptococcus agalactia*, telles qu'elles peuvent être isolées à partir de SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345.

De plus, les séquences nucléotidiques isolées et/ caractérisées en ce qu'elles comprennent une séquence nucléotidique choisie parmi :

- a) une séquence nucléotidique codant pour un polypeptide choisi parmi les séquences SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481;
- b) une séquence nucléotidique comportant au moins 75 %, 80 %, 85 %, 90 %, 95 % ou 98 % d'identité avec une séquence nucléotidique codant pour un polypeptide choisi parmi les séquences SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481;



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- c) une séquence nucléotidique s'hybridant dans des conditions de forte stringence avec une séquence nucléotidique codant pour un polypeptide, choisi parmi les séquences SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481, et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides;
- d) une séquence nucléotidique complémentaire ou d'ARN correspondant à une séquence telle que définie en a), b) ou c);
- e) une séquence nucléotidique d'un fragment représentatif d'une séquence telle que définie en a), b), c) ou d) et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides; et
- f) une séquence telle que définie en a), b), c), d) ou e) modifiée, de préférence comportant au plus 10 %, 5 %, 1 % ou 0,5 % de nucléotides modifiés par rapport à la séquence de référence,
- 15 sont également des objets de l'invention.

Selon une réalisation avantageuse, l'invention a pour objet les séquences nucléotidiques isolées et/ caractérisées en ce qu'elles comprennent une séquence nucléotidique choisie parmi :

- a) une séquence nucléotidique SEQ ID No. 4482 à SEQ ID No. 6617;
- b) une séquence nucléotidique comportant au moins 75 %, 80 %, 85 %, 90 %, 95 % ou 98 % d'identité avec une séquence nucléotidique choisi parmi les séquences SEQ ID No. 2346 à SEQ ID No. 4481;
  - c) une séquence nucléotidique s'hybridant dans des conditions de forte stringence avec une séquence nucléotidique choisi parmi les séquences SEQ ID No. 4482 à SEQ ID No. 6617, et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides;
  - d) une séquence nucléotidique complémentaire ou d'ARN correspondant à une séquence telle que définie en a), b) ou c);
- e) une séquence nucléotidique d'un fragment représentatif d'une séquence telle que définie en a), b), c) ou d) et comprenant au moins 20 nucléotides, de préférence 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 500, 750, 1000 ou 1500 nucléotides; et

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f) une séquence telle que définie en a), b), c), d) ou e) modifiée, de préférence comportant au plus 10 %, 5 %, 1 % ou 0,5 % de nucléotides modifiés par rapport à la séquence de référence,

Par acide nucléique, séquence nucléique ou d'acide nucléique, polynucléotide, oligonucléotide, séquence de polynucléotide, séquence nucléotidique, termes qui seront employés indifféremment dans la présente description, on entend désigner un enchaînement précis de nucléotides, modifiés ou non, permettant de définir un fragment ou une région d'un acide nucléique, comportant ou non des nucléotides non naturels, et pouvant correspondre aussi bien à un ADN double brin, un ADN simple brin qu'à des produits de transcription desdits ADNs. Ainsi, les séquences nucléiques selon l'invention englobent également les PNA (Peptid Nucleic Acid).

Il doit être compris que la présente invention ne concerne pas les séquences nucléotidiques dans leur environnement chromosomique naturel, c'est-à-dire à l'état naturel. Il s'agit de séquences qui ont été isolées et/ou purifiées, c'est-à-dire qu'elles ont été prélevées directement ou indirectement, par exemple par copie, leur environnement ayant été au moins partiellement modifié. On entend ainsi également désigner les acides nucléiques obtenus par synthèse chimique.

Par « pourcentage d'identité » entre deux séquences d'acides nucléiques ou d'acides aminés au sens de la présente invention, on entend désigner un pourcentage de nucléotides ou de résidus d'acides aminés identiques entre les deux séquences à comparer, obtenu après le meilleur alignement, ce pourcentage étant purement statistique et les différences entre les deux séquences étant réparties au hasard et sur toute leur longueur. On entend désigner par "meilleur alignement" ou "alignement optimal", l'alignement pour lequel le pourcentage d'identité déterminé comme ci-après est le plus élevé. Les comparaisons de séquences entre deux séquences d'acides nucléiques ou d'acides aminés sont traditionnellement réalisées en comparant ces séquences après les avoir alignées de manière optimale, ladite comparaison étant réalisée par segment ou par « fenêtre de comparaison » pour identifier et comparer les régions locales de similarité de séquence. L'alignement optimal des séquences pour la comparaison peut être réalisé, outre manuellement, au moyen de l'algorithme d'homologie locale de Smith et Waterman (1981, Ad. App. Math. 2:482), au moyen de l'algorithme d'homologie locale de Neddleman et Wunsch (1970, J. Mol. Biol. 48:443), au moyen de la méthode de recherche de similarité de Pearson et Lipman (1988, Proc.



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Natl. Acad. Sci. USA 85:2444), au moyen de logiciels informatiques utilisant ces algorithmes (GAP, BESTFIT, BLAST P, BLAST N, FASTA et TFASTA dans le Wisconsin Genetics Software Package, Genetics Computer Group, 575 Science Dr., Madison, WI). Afin d'obtenir l'alignement optimal, on utilise de préférence le programme BLAST, avec la matrice BLOSUM 62. On peut également utiliser les matrices PAM ou PAM250.

Le pourcentage d'identité entre deux séquences d'acides nucléiques ou d'acides aminés est déterminé en comparant ces deux séquences alignées de manière optimale, la séquence d'acides nucléiques ou d'acides aminés à comparer pouvant comprendre des additions ou des délétions par rapport à la séquence de référence pour un alignement optimal entre ces deux séquences. Le pourcentage d'identité est calculé en déterminant le nombre de positions identiques pour lesquelles le nucléotide ou le résidu d'acide aminé est identique dans les deux séquences, en divisant ce nombre de positions identiques par le nombre total de positions comparées et en multipliant le résultat obtenu par 100 pour obtenir le pourcentage d'identité entre ces deux séquences.

Par séquences nucléiques présentant un pourcentage d'identité d'au moins 75 %, de préférence 80 %, 85 % ou 90 %, de façon plus préférée 95 % voire 98 %, après alignement optimal avec une séquence de référence, on entend désigner les séquences nucléiques présentant, par rapport à la séquence nucléique de référence, certaines modifications comme en particulier une délétion, une troncation, un allongement, une fusion chimérique et/ou une substitution, notamment ponctuelle, et dont la séquence nucléique présente au moins 75 %, de préférence 80 %, 85 %, 90 %, 95 % ou 98 %, d'identité après alignement optimal avec la séquence nucléique de référence. Il s'agit de préférence de séquences dont les séquences complémentaires sont susceptibles de s'hybrider spécifiquement avec les séquences de référence. De préférence, les conditions d'hybridation spécifiques ou de forte stringence seront telles qu'elles assurent au moins 75 %, de préférence 80 %, 85 %, 90 %, 95 % ou 98 % d'identité après alignement optimal entre l'une des deux séquences et sa séquence complémentaire.

Une hybridation dans des conditions de forte stringence signifie que les conditions de température et de force ionique sont choisies de telle manière qu'elles permettent le maintien de l'hybridation entre deux fragments d'ADN complémentaires. A titre illustratif, des conditions de forte stringence de l'étape d'hybridation aux fins de

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définir les fragments polynucléotidiques décrits ci-dessus, sont avantageusement les suivantes.

L'hybridation ADN-ADN ou ADN-ARN est réalisée en deux étapes : (1) préhybridation à 42°C pendant 3 heures en tampon phosphate (20 mM, pH 7,5) contenant 5 x SSC (1 x SSC correspond à une solution 0,15 M NaCl + 0,015 M citrate de sodium), 50 % de formamide, 7 % de sodium dodécyl sulfate (SDS), 10 x Denhardt's, 5 % de dextran sulfate et 1 % d'ADN de sperme de saumon; (2) hybridation proprement dite pendant 20 heures à une température dépendant de la taille de la sonde (i.e. : 42°C, pour une sonde de taille > 100 nucléotides) suivie de 2 lavages de 20 minutes à 20°C en 2 x SSC + 2 % SDS, 1 lavage de 20 minutes à 20°C en 0,1 x SSC + 0,1 % SDS. Le dernier lavage est pratiqué en 0,1 x SSC + 0,1 % SDS pendant 30 minutes à 60°C pour une sonde de taille > 100 nucléotides. Les conditions d'hybridation de forte stringence décrites ci-dessus pour un polynucléotide de taille plus grande ou plus petite, selon l'enseignement de Sambrook et al., (1989, Molecular cloning : a laboratory manual. 2<sup>nd</sup> Ed. Cold Spring Harbor).

De plus, par fragment représentatif de séquences selon l'invention, on entend désigner tout fragment nucléotidique présentant au moins 15 nucléotides, de préférence au moins 25, 30, 35, 40, 50, 75, 100, 150, 200, 250, 300, 400, 450, 500, 750, 1000 ou 1500 consécutifs de la séquence dont il est issu.

Par fragment représentatif, on entend en particulier une séquence nucléique codant pour un fragment biologiquement actif d'un polypeptide, tel que défini plus loin.

Par fragment représentatif, on entend également les séquences intergéniques, et en particulier les séquences nucléotidiques portant les signaux de régulation (promoteurs, terminateurs, voire enhancers, ...).

Parmi lesdits fragments représentatifs, on préfère ceux ayant des séquences nucléotidiques correspondant à des cadres ouverts de lecture, dénommés séquences ORFs (ORF pour « Open Reading Frame »), compris en général entre un codon d'initiation et un codon stop, ou entre deux codons stop, et codant pour des polypeptides, de préférence d'au moins 100 acides aminés, tel que par exemple, sans s'y limiter, les séquences ORFs qui seront décrites par la suite.

La numérotation des séquences nucléotidiques ORFs qui sera utilisée par la suite dans la présente description correspond à la numérotation des séquences d'acides

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aminés des protéines codées par lesdites ORFs pour les peptides de séquence SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No.2346 à SEQ ID No.4481.

Les fragments représentatifs selon l'invention peuvent être obtenus par exemple par amplification spécifique telle que la PCR ou après digestion par des enzymes de restriction appropriés de séquences nucléotidiques selon l'invention, cette méthode étant décrite en particulier dans l'ouvrage de Sambrook et al.. Les dits fragments représentatifs peuvent également être obtenus par synthèse chimique lorsque leur taille n'est pas trop importante, selon des méthodes bien connues de l'homme du métier.

Parmi les séquences contenant des séquences de l'invention, ou des fragments représentatifs, on entend également les séquences qui sont naturellement encadrées par des séquences qui présentent au moins 75 %, 80 %, 85 %, 90 %, 95 % ou 98 % d'identité avec les séquences selon l'invention.

Par séquence nucléotidique modifiée, on entend toute séquence nucléotidique obtenue par mutagénèse selon des techniques bien connues de l'homme du métier, et comportant des modifications par rapport aux séquences normales, par exemple des mutations dans les séquences régulatrices et/ou promotrices de l'expression du polypeptide, notamment conduisant à une modification du taux d'expression ou de l'activité dudit polypeptide.

Par séquence nucléotidique modifiée, on entend également toute séquence nucléotidique codant pour un polypeptide modifié tel que définit ci-après.

Concernant les séquences nucléiques ou ORF codant pour les peptides de séquence SEQ ID No. 2346 à SEQ ID No. 4481, ces séquences nucléiques ou ORF sont représentées respectivement par les séquences SEQ ID No. 4482 à SEQ ID No. 6617.

L'invention concerne avantageusement une séquence nucléotidique isolée de Streptococcus agalactiae, caractérisée en ce qu'elle est choisie parmi :

- a) une séquence choisie parmi les séquences SEQ ID N° 6194,6236,5497,5791,5103,4705,5610,5234,4926,6331,6247,5842,5741,4921,5090, 5180,4706,4708,5677,6246,6411,5578,6446,6447,5607,6209,6215,5406,5658,4965, de préférence parmi les séquences SEQ ID N°4926,6331,5491,5234,6246,5842;
- b) une séquence nucléotidique comportant au moins 75 % d'identité avec une séquence nucléotidique du a);
- c) une séquence nucléotidique s'hybridant dans des conditions de forte stringence avec une séquence nucléotidique du a) ou b) et comprenant au moins 20 nucléotides;

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- d) une séquence nucléotidique complémentaire ou d'ARN correspondant à une séquence telle que définie en a), b) ou c);
- e) une séquence nucléotidique d'un fragment représentatif d'une séquence telle que définie en a), b), c) ou d) et comprenant au moins 20 nucléotides ; et
- f) une séquence telle que définie en a), b), c), d) ou e) modifiée et comportant au plus 10 % de nucléotides modifiés par rapport à la séquence de référence ;

et en ce qu'elle code pour une protéine de surface avec un motif d'ancrage LPXTG.

L'invention concerne également les polypeptides codés par ces séquences.

L'invention concerne également avantageusement une séquence nucléotidique isolée de *Streptococcus agalactiae* caractérisée en ce qu'elle est choisie parmi les séquences SEQ ID N°6035,6137,6335,6377,6386,4495,4596,4636,4730,4816,4836,4906,4920,4925,5158, 5247, 5306,5417,5450,5486,5559,5591,5677,5732,5799,5800,5861,5923; et en ce qu'elle code pour une lipoprotéine. L'invention concerne également les polypeptides codés par ces séquences.

L'invention concerne également avantageusement une séquence nucléotidique isolée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi les séquences

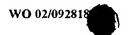
SEQ

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N°4861,6214,6061,6517,6518,6519,4743,6343,6342,5326,4952,5619,5618,5617,5616, 5615,5614,5613,5611,5696,5971,5233,5602,5156,5574,5573,5654,5656,5526,5527,552 9,5534,5625,5626,6223,6229,6230,6231,6232,6233,5764,6095,5089,5466,5465; et en ce qu'elle code pour une protéine impliquée dans la biosynthèse de composés polysaccharidiques de paroi. L'invention concerne également les polypeptides codés par ces séquences.

Les fragments représentatifs selon l'invention peuvent également être des sondes ou amorces, qui peuvent être utilisées dans des procédés de détection, d'identification, de dosage ou d'amplification de séquences nucléiques.

Une sonde ou amorce se définit, au sens de l'invention, comme étant un fragment d'acides nucléiques simple brin ou un fragment double brin dénaturé comprenant par exemple de 12 bases à quelques kb, notamment de 15 à quelques centaines de bases, de préférence de 15 à 50 ou 100 bases, et possédant une spécificité d'hybridation dans des conditions déterminées pour former un complexe d'hybridation avec un acide nucléique cible.



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Les sondes et amorces selon l'invention peuvent être marquées directement ou indirectement par un composé radioactif ou non radioactif par des méthodes bien connues de l'homme du métier, afin d'obtenir un signal détectable et/ou quantifiable (brevet FR 78 10975 et bDNA de Chiron EP 225 807 et EP 510 085).

Les séquences non marquées de polynucléotides selon l'invention peuvent être utilisées directement comme sonde ou amorce.

Les séquences sont généralement marquées pour obtenir des séquences utilisables pour de nombreuses applications. Le marquage des amorces ou des sondes selon l'invention est réalisé par des éléments radioactifs ou par des molécules non radioactives.

Parmi les isotopes radioactifs utilisés, on peut citer le <sup>32</sup>P, le <sup>33</sup>P, le <sup>35</sup>S, le <sup>3</sup>H ou le <sup>125</sup>I. Les entités non radioactives sont sélectionnées parmi les ligands tels la biotine, l'avidine, la streptavidine, la dioxygénine, les haptènes, les colorants, les agents luminescents tels que les agents radioluminescents, chémoluminescents, bioluminescents, fluorescents, phosphorescents.

Les polynucléotides selon l'invention peuvent ainsi être utilisés comme amorce et/ou sonde dans des procédés mettant en oeuvre notamment la technique de PCR (amplification en chaîne par polymérase) (Rolfs et al., 1991, Berlin: Springer-Verlag). Cette technique nécessite le choix de paires d'amorces oligonucléotidiques encadrant le fragment qui doit être amplifié. On peut, par exemple, se référer à la technique décrite dans le brevet américain U.S. N° 4,683,202. Les fragments amplifiés peuvent être identifiés, par exemple après une électrophorèse en gel d'agarose ou de polyacrylamide, ou après une technique chromatographique comme la filtration sur gel ou la chromatographie échangeuse d'ions, puis séquencés. La spécificité de l'amplification peut être contrôlée en utilisant les séquences nucléotidiques de polynucléotides de l'invention comme matrice, des plasmides contenant ces séquences ou encore les produits d'amplification dérivés. Les fragments nucléotidiques amplifiés peuvent être utilisés comme réactifs dans des réactions d'hybridation afin de mettre en évidence la présence, dans un échantillon biologique, d'un acide nucléique cible de séquence complémentaire à celle desdits fragments nucléotidiques amplifiés.

L'invention vise également les acides nucléiques susceptibles d'être obtenus par amplification à l'aide d'amorces selon l'invention.

D'autres techniques d'amplification de l'acide nucléique cible peuvent être avantageusement employées comme alternative à la PCR (PCR-like) à l'aide de couple

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d'amorces de séquences nucléotidiques selon l'invention. Par PCR-like on entend désigner toutes les méthodes mettant en œuvre des reproductions directes ou indirectes des séquences d'acides nucléiques, ou bien dans lesquelles les systèmes de marquage ont été amplifiés, ces techniques sont bien entendu connues. En général il s'agit de l'amplification de l'ADN par une polymérase ; lorsque l'échantillon d'origine est un ARN il convient préalablement d'effectuer une transcription reverse. Il existe actuellement de très nombreux procédés permettant cette amplification, comme par exemple la technique SDA (Strand Displacement Amplification) ou technique d'amplification à déplacement de brin (Walker et al., 1992, Nucleic Acids Res. 20:1691), la technique TAS (Transcription-based Amplification System) décrite par Kwoh et al. (1989, Proc. Natl. Acad. Sci., USA, 86, 1173), la technique 3SR (Self-Sustained Sequence Replication) décrite par Guatelli et al. (1990, Proc. Natl. Acad. Sci., USA 87:1874), la technique NASBA (Nucleic Acid Sequence Based Amplification) décrite par Kievitis et al. (1991, J. Virol. Methods, 35, 273), la technique TMA (Transcription Mediated Amplification), la technique LCR (Ligase Chain Reaction) décrite par Landegren et al. (1988, Science 241, 1077), la technique de RCR (Repair Chain Reaction) décrite par Segev (1992, Kessler C. Springer Verlag, Berlin, New-York, 197-205), la technique CPR (Cycling Probe Reaction) décrite par Duck et al. (1990, Biotechniques, 9, 142), la technique d'amplification à la Q-béta-réplicase décrite par Miele et al. (1983, J. Mol. Biol., 171, 281). Certaines de ces techniques ont depuis été perfectionnées.

Dans le cas où le polynucléotide cible à détecter est un ARNm, on utilise avantageusement, préalablement à la mise en œuvre d'une réaction d'amplification à l'aide des amorces selon l'invention ou à la mise en œuvre d'un procédé de détection à l'aide des sondes de l'invention, une enzyme de type transcriptase inverse afin d'obtenir un ADNc à partir de l'ARNm contenu dans l'échantillon biologique. L'ADNc obtenu servira alors de cible pour les amorces ou les sondes mises en oeuvre dans le procédé d'amplification ou de détection selon l'invention.

La technique d'hybridation de sondes peut être réalisée de manières diverses (Matthews et al., 1988, Anal. Biochem., 169, 1-25). La méthode la plus générale consiste à immobiliser l'acide nucléique extrait des cellules de différents tissus ou de cellules en culture sur un support (tels que la nitrocellulose, le nylon, le polystyrène) et à incuber, dans des conditions bien définies, l'acide nucléique cible immobilisé avec la sonde. Après l'hybridation, l'excès de sonde est éliminé et les molécules hybrides

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formées sont détectées par la méthode appropriée (mesure de la radioactivité, de la fluorescence ou de l'activité enzymatique liée à la sonde).

Selon un autre mode de mise en œuvre des sondes nucléiques selon l'invention, ces dernières peuvent être utilisées comme sondes de capture. Dans ce cas, une sonde, dite « sonde de capture », est immobilisée sur un support et sert à capturer par hybridation spécifique l'acide nucléique cible obtenu à partir de l'échantillon biologique à tester et l'acide nucléique cible est ensuite détecté grâce à une seconde sonde, dite « sonde de détection », marquée par un élément facilement détectable.

Parmi les fragments d'acides nucléiques intéressants, il faut ainsi citer en particulier les oligonucléotides anti-sens, c'est-à-dire dont la structure assure, par hybridation avec la séquence cible, une inhibition de l'expression du produit correspondant. Il faut également citer les oligonucléotides sens qui, par interaction avec des protéines impliquées dans la régulation de l'expression du produit correspondant, induiront soit une inhibition, soit une activation de cette expression.

De façon préférée, les sondes ou amorces selon l'invention sont immobilisées sur un support, de manière covalente ou non covalente. En particulier, le support peut être une puce à ADN ou un filtre à haute ou moyenne densité, également objets de la présente invention (brevets WO 97/29212, WO 98/27317, WO 97/10365 et WO 92/10588).

On entend désigner par puce à ADN ou filtre haute densité, un support sur lequel sont fixées des séquences d'ADN, chacune d'entre elles pouvant être repérée par sa localisation géographique. Ces puces ou filtres diffèrent principalement par leur taille, le matériau du support, et éventuellement le nombre de séquences d'ADN qui y sont fixées.

On peut fixer les sondes ou amorces selon la première invention sur des supports solides, en particulier les puces à ADN, par différents procédés de fabrication. En particulier, on peut effectuer une synthèse in situ par adressage photochimique ou par jet d'encre. D'autres techniques consistent à effectuer une synthèse ex situ et à fixer les sondes sur le support de la puce à ADN par adressage mécanique, électronique ou par jet d'encre. Ces différents procédés sont bien connus de l'homme du métier.

Une séquence nucléotidique (sonde ou amorce) selon l'invention permet donc la détection et/ou l'amplification de séquences nucléiques spécifiques. En particulier, la détection de cesdites séquences est facilitée lorsque la sonde est fixée sur une puce à ADN, ou à un filtre haute densité.

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L'utilisation de puces à ADN ou de filtres à haute densité permet en effet de déterminer l'expression de gènes dans un organisme présentant une séquence génomique proche de Streptococcus agalactiae et le typage de la souche en cause.

La séquence génomique de *Streptococcus agalactiae*, complétée par l'identification des gènes de ces organismes, telle que présentée dans la présente invention, sert de base à la construction de ces puces à ADN ou filtre.

La préparation de ces filtres ou puces consiste à synthétiser des oligonucléotides, correspondant aux extrémités 5' et 3' des gènes ou à des fragments plus internes pour amplifier des fragments d'une taille adaptée, par exemple comprise environ entre 300 et 800 bases. Ces oligonucléotides sont choisis en utilisant la séquence génomique et ses annotations divulguées par la présente invention. La température d'appariement des ces oligonucléotides aux places correspondantes sur l'ADN doit être approximativement la même pour chaque oligonucleotide. Ceci permet de préparer des fragments d'ADN correspondant à chaque gène par l'utilisation de conditions de PCR appropriées dans un environnement hautement automatisé. Les fragments amplifiés sont ensuite immobilisés sur des filtres ou des supports en verre, silicium ou polymères synthétiques et ces milieux sont utilisés pour l'hybridation.

La disponibilité de tels filtres et/ou puces et de la séquence génomique correspondante annotée permet d'étudier l'expression de grands ensembles, voire de la totalité des gènes dans les micro-organismes associés à *Streptococcus agalactiae* et *Streptococcus agalactiae* CIP 82.45 (ATCC 12403), en préparant les ADNs complémentaires, et en les hybridant à l'ADN ou aux oligonucléotides immobilisés sur les filtres ou les puces. De même, les filtres et/ou les puces permettent d'étudier la variabilité des souches ou des espèces, en préparant l'ADN de ces organismes et en les hybridant à l'ADN ou aux oligonucléotides immobilisés sur les filtres ou les puces.

Les différences entre les séquences génomiques des différentes souches ou espèces peuvent grandement affecter l'intensité de l'hybridation et, par conséquent, perturber l'interprétation des résultats. Il peut donc être nécessaire d'avoir la séquence précise des gènes de la souche que l'on souhaite étudier. La méthode de détection des gènes décrite plus loin en détail, impliquant la détermination de la séquence de fragments aléatoires d'un génome, et les organisant d'après la séquence du génome de *Streptococcus agalactiae*, notamment de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) divulguée dans la présente invention, peut être très utile.



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Les séquences nucléotidiques selon l'invention peuvent être utilisées dans des puces à ADN pour effectuer l'analyse de mutations. Cette analyse repose sur la constitution de puces capables d'analyser chaque base d'une séquence nucléotidique selon l'invention. On pourra notamment à cette fin mettre en œuvre les techniques de microséquençage sur puce à ADN. Les mutations sont détectées par extension d'amorces immobilisées hybridant à la matrice des séquences analysées, juste en position adjacente de celle du nucléotide muté recherché. Une matrice simple brin, ARN ou ADN, des séquences à analyser sera avantageusement préparée selon des méthodes classiques, à partir de produits amplifiés selon les techniques de type PCR. Les matrices d'ADN simple brin, ou d'ARN ainsi obtenues sont alors déposées sur la puce à ADN, dans des conditions permettant leur hybridation spécifique aux amorces immobilisées. Une polymérase thermostable, par exemple la Tth ou la Taq ADN polymérase, étend spécifiquement l'extrémité 3' de l'amorce immobilisée avec un analogue de nucléotide marqué complémentaire du nucléotide en position du site variable ; par exemple, un cyclage thermique est réalisé en présence des didéoxyribonucléotides fluorescents. Les conditions expérimentales seront adaptées notamment aux puces employées, aux amorces immobilisées, aux polymérases employées, et au système de marquage choisi. Un avantage du microséquençage, par rapport aux techniques basées sur l'hybridation de sondes, est qu'il permet d'identifier tous les nucléotides variables avec une discrimination optimale dans des conditions de réactions homogènes ; utilisé sur des puces à ADN, il permet une résolution et une spécificité optimales pour la détection routinière et industrielle de mutations en multiplex.

Une puce à ADN ou un filtre peut être un outil extrêmement intéressant pour la détermination, la détection et/ou l'identification d'un micro-organisme. Ainsi, on préfère également les puces à ADN selon l'invention qui contiennent en outre au moins une séquence nucléotidique d'un micro-organisme autre que *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou *Streptococcus agalactiae*, immobilisée sur le support de ladite puce. De préférence, le micro-organisme choisi l'est parmi les bactéries du genre *Streptococcus* (ci-après désignées comme bactéries associées à *Streptococcus agalactiae*), ou les variants de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403).

Une puce à ADN ou un filtre selon l'invention est un élément très utile de certains kits ou nécessaires pour la détection et/ou l'identification de micro-organismes, en particulier les bactéries appartenant à l'espèce Streptococcus agalactiae ou les micro-organismes associés, également objets de l'invention.

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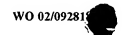
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Par ailleurs, les puces à ADN ou les filtres selon l'invention, contenant des sondes ou amorces spécifiques de Streptococcus agalactiae, sont des éléments très avantageux de kits ou nécessaires pour la détection et/ou la quantification de l'expression de gènes de Streptococcus agalactiae (ou de micro-organismes associés).

En effet, le contrôle de l'expression des gènes est un point critique pour optimiser la croissance et le rendement d'une souche, soit en permettant l'expression d'un ou plusieurs gènes nouveaux, soit en modifiant l'expression de gènes déjà présents dans la cellule. La présente invention fournit l'ensemble des séquences naturellement actives chez Streptococcus agalactiae permettant l'expression des gènes. Elle permet ainsi la détermination de l'ensemble des séquences exprimées chez Streptococcus agalactiae. Elle fournit également un outil permettant de repérer les gènes dont l'expression suit un schéma donné. Pour réaliser cela, l'ADN de tout ou partie des gènes de Streptococcus agalactiae peut être amplifié grâce à des amorces selon l'invention, puis fixé à un support comme par exemple le verre ou le nylon ou une puce à ADN, afin de construire un outil permettant de suivre le profil d'expression de ces gènes. Cet outil, constitué de ce support contenant les séquences codantes sert de matrice d'hybridation à un mélange de molécules marquées reflétant les ARNs messagers exprimés dans la cellule (en particulier les sondes marquées selon l'invention). En répétant cette expérience à différents instants et en combinant l'ensemble de ces données par un traitement approprié, on obtient alors les profils d'expression de l'ensemble de ces gènes. La connaissance des séquences qui suivent un schéma de régulation donné peut aussi être mise à profit pour rechercher de manière dirigée, par exemple par homologie, d'autres séquences suivant globalement, mais de manière légèrement différente le même schéma de régulation. En complément, il est possible d'isoler chaque séquence de contrôle présente en amont des segments servant de sondes et d'en suivre l'activité à l'aide de moyen approprié comme un gène rapporteur (luciférase, β-galactosidase, GFP). Ces séquences isolées peuvent ensuite être modifiées et assemblées par ingénierie métabolique avec des séquences d'intérêt en vue de leur expression optimale.

L'invention concerne également les polypeptides codés par une séquence nucléotidique selon l'invention, de préférence, par un fragment représentatif des séquences précédentes et correspondant à une séquence ORF. En particulier, les polypeptides de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) de SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481 sont objet de l'invention.



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L'invention comprend également les polypeptides caractérisés en ce qu'ils comprennent un polypeptide choisi parmi :

- a) un polypeptide selon l'invention;
- b) un polypeptide présentant au moins 80 % de préférence 85 %, 90 %, 95 % et 98 % d'identité avec un polypeptide selon l'invention;
- c) un fragment d'au moins 5 acides aminés, de préférence d'au moins 10, 15, 20, 25, 30, 40, 50, 75 et 100 acides aminés d'un polypeptide selon l'invention, ou tel que défini en b);
- d) un fragment biologiquement actif d'un polypeptide selon l'invention, ou tel que défini en b) ou c) ; et
  - e) un polypeptide selon l'invention, ou tel que défini en b), c) ou d) modifié et comportant au plus 10 %, 5 % ou 1 % d'acides aminés modifiés par rapport à la séquence de référence.

Les séquences nucléotidiques codant pour les polypeptides décrits précédemment sont également objet de l'invention.

Dans la présente description, les termes polypeptides, séquences polypeptidiques, peptides et protéines sont interchangeables. Le terme polypeptide comprend toute séquence d'acides aminés permettant de générer une réponse anticorps.

Il doit être compris que l'invention ne concerne pas les polypeptides sous forme naturelle, c'est-à-dire qu'ils ne sont pas pris dans leur environnement naturel. En revanche, elle concerne ceux qui ont pu être isolés ou obtenus par purification à partir de sources naturelles, ou bien obtenus par recombinaison génétique, ou par synthèse chimique, et qu'ils peuvent alors comporter des acides aminés non naturels comme cela sera décrit plus loin.

Par polypeptide présentant un certain pourcentage d'identité avec un autre, que l'on désignera également par polypeptide homologue, on entend désigner les polypeptides présentant par rapport aux polypeptides naturels, certaines modifications, en particulier une délétion, addition ou substitution d'au moins un acide aminé, une troncation, un allongement, une solution chimérique et/ou une mutation, ou les polypeptides présentant des modifications post-traductionnelles. Parmi les polypeptides homologues, on préfère ceux dont la séquence d'acides aminés présentent au moins 80 %, de préférence 85 %, 90 %, 95 % et 98 % d'homologie avec les séquences d'acides aminés des polypeptides selon l'invention. Dans le cas d'une substitution, un ou plusieurs acide(s) aminé(s) consécutif(s) ou non consécutif(s) sont remplacés par des

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acides aminés « équivalents ». L'expression « acides aminés équivalents » vise ici à désigner tout acide aminé susceptible d'être substitué à l'un des acides aminés de la structure de base sans cependant modifier essentiellement les activités biologiques des peptides correspondant telles qu'elles seront définies par la suite.

Ces acides aminés équivalents peuvent être déterminés soit en s'appuyant sur leur homologie de structure avec les acides aminés auxquels ils se substituent, soit sur des résultats d'essais comparatifs d'activité biologique entre les différents polypeptides susceptibles d'être effectués.

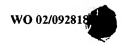
A titre d'exemple, on mentionne les possibilités de substitution susceptibles d'être effectuées sans qu'il résulte en une modification approfondie de l'activité biologique du polypeptide modifié correspondant. On peut remplacer ainsi la leucine par la valine ou l'isoleucine, l'acide aspartique par l'acide glutamine, la glutamine par l'asparagine, l'arginine par la lysine, etc., les substitutions inverses étant naturellement envisageables dans les mêmes conditions.

Les polypeptides homologues correspondent également aux polypeptides codés par les séquences nucléotidiques homologues ou identiques, telles que définies précédemment et comprennent ainsi dans la présente définition des polypeptides mutés ou correspondant à des variations inter ou intra espèces, pouvant exister chez *Streptococcus*, et qui correspondent notamment à des troncatures, substitutions, délétions et/ou additions, d'au moins un résidu d'acides aminés.

Il est entendu que l'on calcule le pourcentage d'identité entre deux polypeptides de la même façon qu'entre deux séquences d'acides nucléiques. Ainsi, le pourcentage d'identité entre deux polypeptides est calculé après alignement optimal de ces deux séquences, sur une fenêtre d'homologie maximale. Pour définir ladite fenêtre d'homologie maximale, on peut utiliser les mêmes algorithmes que pour les séquences d'acide nucléique.

Par fragment biologiquement actif d'un polypeptide selon l'invention, on entend désigner en particulier un fragment de polypeptide, tel que défini ci-après, présentant au moins une des caractéristiques biologiques des polypeptides selon l'invention, notamment en ce qu'il est capable d'exercer de manière générale une activité même partielle, telle que par exemple :

- une activité enzymatique (métabolique) ou une activité pouvant être impliquée dans la biosynthèse ou la biodégradation de composés organiques ou inorganiques;



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- une activité structurelle (enveloppe cellulaire, molécule chaperonne, ribosome);
  - une activité de transport (d'énergie, d'ion) ; ou dans la sécrétion de protéine ;
- une activité dans le processus de réplication, amplification, préparation, transcription, traduction ou maturation, notamment de l'ADN, de l'ARN ou des protéines.

Par fragment de polypeptide selon l'invention, on entend désigner un polypeptide comportant au minimum 5 acides aminés, de préférence d'au moins 10, 15, 20, 25, 30, 40, 50, 75, 100 et 150 acides aminés.

Les fragments de polypeptides peuvent correspondre à des fragments isolés ou purifiés naturellement présents dans les souches de *Streptococcus*, ou à des fragments qui peuvent être obtenus par clivage dudit polypeptide par une enzyme protéolitique telle que la trypsine ou la chymotrypsine ou la collagénase, par un réactif chimique (bromure de cyanogène, CNBr) ou en plaçant ledit polypeptide dans un environnement très acide (par exemple à pH = 2,5). Des fragments polypeptidiques peuvent également être préparés par synthèse chimique, à partir d'hôtes transformés par un vecteur d'expression selon l'invention qui contiennent un acide nucléique permettant l'expression dudit fragment, et placé sous le contrôle des éléments de régulation et/ou d'expression appropriés.

Par « polypeptide modifié » d'un polypeptide selon l'invention, on entend désigner un polypeptide obtenu par recombinaison génétique ou par synthèse chimique comme décrit plus loin, qui présente au moins une modification par rapport à la séquence normale. Ces modifications peuvent être notamment portées sur des acides aminés nécessaires pour la spécificité ou l'efficacité de l'activité, ou à l'origine de la conformation structurale, de la charge, ou de l'hydrophobicité du polypeptide selon l'invention. On peut ainsi créer des polypeptides d'activité équivalente, augmentée ou diminuée, ou de spécificité équivalente, plus étroite ou plus large. Parmi les polypeptides modifiés, il faut citer les polypeptides dans lesquels jusqu'à cinq acides aminés peuvent être modifiés, tronqués à l'extrémité N ou C-terminale, ou bien délétés, ou ajoutés.

Comme cela est indiqué, les modifications d'un polypeptide ont pour objectif notamment :

- de permettre sa mise en œuvre dans des procédés de biosynthèse ou de biodégradation de composés organiques ou inorganiques,

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- de permettre sa mise en œuvre dans des procédés de réplication, d'amplification, de réparation et règle de transcription, de traduction, ou de maturation notamment de l'ADN, l'ARN, ou de protéines,
  - de permettre sa sécrétion améliorée,
- de modifier sa solubilité, l'efficacité ou la spécificité de son activité, ou encore de faciliter sa purification.

La synthèse chimique présente également l'avantage de pouvoir utiliser des acides aminés non naturels ou des liaisons non peptidiques. Ainsi, il peut être intéressant d'utiliser des acides aminés non naturels, par exemple sous forme D, ou des analogues d'acides aminés, notamment des formes souffrées.

La présente invention fournit la séquence nucléotidique du génome de Streptococcus agalactiae CIP 82.45 (ATCC 12403) sous forme de contigs, ainsi que certaines séquences polypeptidiques.

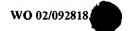
D'une manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la biosynthèse des acides aminés.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la biosynthèse des cofacteurs, groupes prosthétiques et transporteurs.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide d'enveloppe cellulaire ou présent à la surface de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou pour un de ses fragments.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la machinerie cellulaire.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme intermédiaire central.



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De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme énergétique.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme des acides gras et des phospholipides.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme des nucléotides, des purines, des pyrimidines ou nucléosides.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans les fonctions de régulation.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de réplication.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de transcription.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de traduction.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de transport et de liaison des protéines.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus* 

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agalactiae CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans l'adaptation aux conditions atypiques.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments dans la sensibilité aux médicaments et analogues.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans les fonctions relatives aux transposons.

De manière préférée, l'invention est relative à une séquence nucléotidique selon l'invention, caractérisée en ce qu'elle code pour un polypeptide spécifique de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la biosynthèse des acides aminés.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la biosynthèse des cofacteurs, groupes prosthétiques et transporteurs.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide d'enveloppe cellulaire ou de surface de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans la machinerie cellulaire.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme intermédiaire central.

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Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme énergétique.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme des acides gras et des phospholipides.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le métabolisme des nucléotides, des purines, des pyrimidines ou nucléosides.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans les fonctions de régulation.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de réplication.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de transcription.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de traduction.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans le processus de transport et de liaison des protéines.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de

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Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans l'adaptation aux conditions atypiques.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments dans la sensibilité aux médicaments et analogues.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments impliqué dans les fonctions relatives aux transposons.

Sous un autre aspect, de manière préférée, l'invention a pour objet un polypeptide selon l'invention, caractérisé en ce qu'il s'agit d'un polypeptide spécifique de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un de ses fragments.

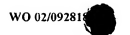
L'invention a également pour objet les opérons impliqués dans la synthèse d'antibiotiques et/ou de toxines.

Le tableau 1 fournit la liste de certains polypeptides selon l'invention, ainsi que leur localisation dans les séquences SEQ ID No. 1 à SEQ ID No. 139, et les analogies observées après comparaison dans les bases de données.

Il est important de noter toutefois qu'un organisme vivant est un tout et doit être pris comme tel. Ainsi, afin de pouvoir se développer et exhiber ses propriétés, tout organisme a besoin d'interactions entre les différentes voies métaboliques. Ainsi, la classification énoncée ci-dessus ne doit pas être considérée comme limitative, un gène pouvant être impliqué dans deux voies métaboliques distinctes.

La présente invention a également pour objet les séquences nucléotidiques et/ou de polypeptides selon l'invention, caractérisées en ce que lesdites séquences sont enregistrées sur un support d'enregistrement dont la forme et la nature facilitent la lecture, l'analyse et/ou l'exploitation de ladite ou desdites séquence(s). Ces supports peuvent également contenir d'autres informations extraites de la présente invention, notamment les analogies avec des séquences déjà connues, et/ou des informations concernant les séquences nucléotidiques et/ou de polypeptides d'autres microorganismes afin de faciliter l'analyse comparative et l'exploitation des résultats obtenus.

Parmi cesdits supports d'enregistrement, on préfère en particulier les supports lisibles par un ordinateur, tels les supports magnétiques, optiques, électriques ou



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hybrides, en particulier les disquettes informatiques, les CD-ROM, les serveurs informatiques. De tels supports d'enregistrement sont également objet de l'invention.

Les supports d'enregistrement selon l'invention, avec les informations apportées, sont très utiles pour le choix d'amorces ou de sondes nucléotidiques pour la détermination de gènes dans *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou souches proches de cet organisme. De même, l'utilisation de ces supports pour l'étude du polymorphisme génétique de souches proches de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403), en particulier par la détermination des régions de colinéarité, est très utile dans la mesure où ces supports fournissent non seulement la séquence nucleotidique du génome de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403), mais egalement l'organisation génomique dans ladite séquence. Ainsi, les utilisations de supports d'enregistrement selon l'invention sont également des objets de l'invention.

L'analyse d'homologie entre différentes séquences s'effectue en effet avantageusement à l'aide de logiciels de comparaison de séquences, tels le logiciel Blast, ou les logiciels de la trousse GCG, décrits précédemment.

L'invention vise également les vecteurs de clonage et/ou d'expression, qui contiennent une séquence nucléotidique selon l'invention.

Les vecteurs selon l'invention comportent de préférence des éléments qui permettent l'expression et/ou la sécrétion des séquences nucléotidiques dans une cellule hôte déterminée.

Le vecteur doit alors comporter un promoteur, des signaux d'initiation et de terminaison de la traduction, ainsi que des régions appropriées de régulation de la transcription. Il doit pouvoir être maintenu de façon stable dans la cellule hôte et peut éventuellement posséder des signaux particuliers qui spécifient la sécrétion de la protéine traduite. Ces différents éléments sont choisis et optimisés par l'homme du métier en fonction de l'hôte cellulaire utilisé. A cet effet, les séquences nucléotidiques selon l'invention peuvent être insérées dans des vecteurs à réplication autonome au sein de l'hôte choisi, ou être des vecteurs intégratifs de l'hôte choisi.

De tels vecteurs sont préparés par des méthodes couramment utilisées par l'homme du métier, et les clones résultant peuvent être introduits dans un hôte approprié par des méthodes standards, telles que la lipofection, l'électroporation, le choc thermique, ou des méthodes chimiques.

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Les vecteurs selon l'invention sont par exemple des vecteurs d'origine plasmidique ou virale. Ils sont utiles pour transformer des cellules hôtes afin de cloner ou d'exprimer les séquences nucléotidiques selon l'invention.

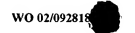
L'invention comprend également les cellules hôtes transformées par un vecteur selon l'invention.

L'hôte cellulaire peut être choisi parmi des systèmes procaryotes ou eucaryotes, par exemple les cellules bactériennes mais également les cellules de levure ou les cellules animales, en particulier les cellules de mammifères. On peut également utiliser des cellules d'insectes ou des cellules de plantes. Les cellules hôtes préférées selon l'invention sont en particulier les cellules procaryotes, de préférence les bactéries appartenant au genre *Streptococcus*, à l'espèce *Streptococcus agalactiae*, plus particulièrement *Streptococcus agalactiae* CIP 82.45 (ATCC 12403), ou les microorganismes associés à l'espèce *Streptococcus agalactiae*.

L'invention concerne également les végétaux et les animaux, excepté l'homme, qui comprennent une cellule transformée selon l'invention. Les cellules transformées selon l'invention sont utilisables dans des procédés de préparation de polypeptides recombinants selon l'invention. Les procédés de préparation d'un polypeptide selon l'invention sous forme recombinante, caractérisés en ce qu'ils mettent en œuvre un vecteur et/ou une cellule transformée par un vecteur selon l'invention sont eux-mêmes compris dans la présente invention. De préférence, on cultive une cellule transformée par un vecteur selon l'invention dans des conditions qui permettent l'expression dudit polypeptide et on récupère ledit peptide recombinant.

Ainsi qu'il a été dit, l'hôte cellulaire peut être choisi parmi des systèmes procaryotes ou eucaryotes. En particulier, il est possible d'identifier des séquences nucléotidiques selon l'invention, facilitant la sécrétion dans un tel système procaryote ou eucaryote. Un vecteur selon l'invention portant une telle séquence peut donc être avantageusement utilisé pour la production de protéines recombinantes, destinées à être sécrétées. En effet, la purification de ces protéines recombinantes d'intérêt sera facilitée par le fait qu'elles sont présentent dans le surnageant de la culture cellulaire plutôt qu'à l'intérieur des cellules hôtes.

On peut également préparer les polypeptides selon l'invention par synthèse chimique. Un tel procédé de préparation est également un objet de l'invention. L'homme du métier connaît les procédés de synthèse chimique, par exemple les techniques mettant en œuvre des phases solides (voir notamment Steward et al., 1984,



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Solid phase peptides synthesis, Pierce Chem. Company, Rockford, 111, 2ème éd., (1984)) ou des techniques utilisant des phases solides partielles, par condensation de fragments ou par une synthèse en solution classique. Les polypeptides obtenus par synthèse chimique et pouvant comporter des acides aminés non naturels correspondants sont également compris dans l'invention.

L'invention est en outre relative à des polypeptides hybrides présentant au moins un polypeptide ou un de ses fragments selon l'invention, et une séquence d'un polypeptide susceptible d'induire une réponse immunitaire chez l'homme ou l'animal.

Avantageusement, le déterminant antigénique est tel qu'il est susceptible d'induire une réponse humorale et/ou cellulaire.

Un tel déterminant pourra comprendre un polypeptide ou un de ses fragments selon l'invention sous forme glycosylée, utilisé en vue d'obtenir des compositions immunogènes susceptibles d'induire la synthèse d'anticorps dirigés contre des épitopes multiples. Les dits polypeptides ou leurs fragments glycosylés font également partie de l'invention.

Ces molécules hybrides peuvent être constituées en partie d'une molécule porteuse de polypeptides ou de leurs fragments selon l'invention, associée à une partie éventuellement immunogène, en particulier un épitope de la toxine diphtérique, la toxine tétanique, un antigène de surface du virus de l'hépatite B (brevet FR 79 21811), l'antigène VP1 du virus de la poliomyélite ou toute autre toxine ou antigène viral ou bactérien.

Les procédés de synthèse des molécules hybrides englobent les méthodes utilisées en génie génétique pour construire des séquences nucléotidiques hybrides codant pour les séquences polypeptidiques recherchées. On pourra, par exemple, se référer avantageusement à la technique d'obtention de gènes codant pour des protéines de fusion décrite par Minton en 1984.

Les dites séquences nucléotidiques hybrides codant pour un polypeptide hybride ainsi que les polypeptides hybrides selon l'invention, caractérisés en ce qu'il s'agit de polypeptides recombinants obtenus par l'expression desdites séquences nucléotidiques hybrides, font également partie de l'invention.

L'invention comprend également les vecteurs caractérisés en ce qu'ils contiennent une desdites séquences nucléotidiques hybrides. Les cellules hôtes transformées par lesdits vecteurs, les animaux transgéniques comprenant une desdites cellules transformées ainsi que les procédés de préparation de polypeptides

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recombinants utilisant lesdits veoteurs, lesdites cellules transformées et/ou lesdits animaux transgéniques font également partie de l'invention.

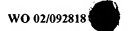
Le couplage entre un polypeptide selon l'invention et un polypeptide immunogène peut être effectué par voie chimique, ou par voie biologique. Ainsi, selon l'invention, il est possible d'introduire un ou plusieurs élément(s) de liaison, notamment des acides aminés pour faciliter les réactions de couplage entre le polypeptide selon l'invention, et le polypeptide immunostimulateur, le couplage covalent de l'antigène immunostimulateur pouvant être réalisé à l'extrémité N ou C-terminale du polypeptide selon l'invention. Les réactifs bifonctionnels permettant ce couplage sont déterminés en fonction de l'extrémité choisie pour réaliser ce couplage, et les techniques de couplage sont bien connues de l'homme du métier.

Les conjugués issus d'un couplage de peptides peuvent être également préparés par recombinaison génétique. Le peptide hybride (conjugué) peut en effet être produit par des techniques d'ADN recombinant, par insertion ou addition à la séquence d'ADN codant pour le polypeptide selon l'invention, d'une séquence codant pour le ou les peptide(s) antigène(s), immunogène(s) ou haptène(s). Ces techniques de préparation de peptides hybrides par recombinaison génétique sont bien connues de l'homme du métier (voir par exemple Makrides, 1996, Microbiological Reviews 60,512-538).

De préférence, ledit polypeptide immunitaire est choisi dans le groupe des peptides contenant les anatoxines, notamment le toxoïde diphtérique ou le toxoïde tétanique, les protéines dérivées du Streptocoque (comme la protéine de liaison à la séralbumine humaine), les protéines membranaires OMPA et les complexes de protéines de membranes externes, les vésicules de membranes externes ou les protéines de chocs thermiques.

Les polypeptides hybrides selon l'invention sont très utiles pour obtenir des anticorps monoclonaux ou polyclonaux, capables de reconnaître spécifiquement les polypeptides selon l'invention. En effet, un polypeptide hybride selon l'invention permet la potentialisation de la réponse immunitaire, contre le polypeptide selon l'invention couplé à la molécule immunogène. De tels anticorps monoclonaux ou polyclonaux, leurs fragments, ou les anticorps chimériques, reconnaissant les polypeptides selon l'invention, sont également objet de l'invention.

Les anticorps monoclonaux spécifiques peuvent être obtenus selon la méthode classique de culture d'hybridome décrite par Köhler et Milstein (1975, Nature 256, 495).



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Les anticorps selon l'invention sont par exemple des anticorps chimériques, des anticorps humanisés, des fragments Fab, ou F(ab')<sup>2</sup>. Ils peuvent également se présenter sous forme d'immunoconjugués ou d'anticorps marqués afin d'obtenir un signal détectable et/ou quantifiable.

Ainsi, les anticorps selon l'invention peuvent être employés dans un procédé pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé dans un échantillon biologique, caractérisé en ce qu'il comprend les étapes suivantes:

- a) mise en contact de l'échantillon biologique avec un anticorps selon l'invention;
- b) mise en évidence du complexe antigène-anticorps éventuellement formé.

Les anticorps selon la présente invention sont également utilisables afin de détecter une expression d'un gène de Streptococcus agalactiae ou de micro-organismes associés. En effet, la présence du produit d'expression d'un gène reconnu par un anticorps spécifique dudit produit d'expression peut être détectée par la présence d'un complexe antigène-anticorps formé après la mise en contact de la souche de Streptococcus agalactiae ou du micro-organisme associé avec un anticorps selon l'invention. La souche bactérienne utilisée peut avoir été « préparée », c'est-à-dire centrifugée, lysée, placée dans un réactif approprié pour la constitution du milieu propice à la réaction immunologique. En particulier, on préfère un procédé de détection de l'expression dans le gène, correspondant à un Western blot, pouvant être effectué après une électrophorèse sur gel de polyacrylamide d'un lysat de la souche bactérienne, en présence ou en l'absence de conditions réductrices (SDS-PAGE). Après migration et séparation des protéines sur le gel de polyacrylamide, on transfère lesdites protéines sur une membrane appropriée (par exemple en nylon) et on détecte la présence de la protéine ou du polypeptide d'intérêt, par mise en contact de ladite membrane avec un anticorps selon l'invention.

Ainsi, la présente invention comprend également les kits ou nécessaires pour la mise en œuvre d'un procédé tel que décrit (de détection de l'expression d'un gène de Streptococcus agalactiae ou d'un micro-organisme associé, ou pour la détection et/ou l'identification de bactéries appartenant à l'espèce Streptococcus agalactiae ou un micro-organisme associé), comprenant les éléments suivants :

- a) un anticorps polyclonal ou monoclonal selon l'invention;
- b) éventuellement, les réactifs pour la constitution du milieu propice à la réaction immunologique;

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c) éventuellement, les réactifs permettant la mise en évidence des complexes antigèneanticorps produits par la réaction immunologique.

Les polypeptides et les anticorps selon l'invention peuvent avantageusement être immobilisés sur un support, notamment une puce à protéines. Une telle puce à protéines est un objet de l'invention, et peut également contenir au moins un polypeptide d'un micro-organisme autre que *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou un anticorps dirigé contre un composé d'un micro-organisme autre que *Streptococcus agalactiae* CIP 82.45 (ATCC 12403).

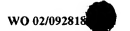
Les puces à protéines ou filtres à haute densité contenant des protéines selon l'invention peuvent être construites de la même manière que les puces à ADN selon l'invention. En pratique, on peut effectuer la synthèse des polypeptides directement fixés sur la puce à protéines, ou effectuer une synthèse ex situ suivie d'une étape de fixation sur ladite puce du polypeptide synthétisé. Cette dernière méthode est préférable, lorsque l'on désire fixer des protéines de taille importante sur le support, ces protéines étant avantageusement préparées par génie génétique. Toutefois, si l'on ne désire fixer que des peptides sur le support de ladite puce, il peut être plus intéressant de procéder à la synthèse desdits peptides directement in situ.

Les puces à protéines selon l'invention peuvent être avantageusement utilisées dans des kits ou nécessaires pour la détection et/ou l'identification de bactéries associées à l'espèce Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou à un microorganisme, ou de façon plus générale dans des kits ou nécessaires pour la détection et/ou l'identification de micro-organismes. Lorsque l'on fixe les polypeptides selon l'invention sur les puces à ADN, on recherche la présence d'anticorps dans les échantillons testés, la fixation d'un anticorps selon l'invention sur le support de la puce à protéines permettant l'identification de la protéine dont ledit anticorps est spécifique.

De préférence, on fixe un anticorps selon l'invention sur le support de la puce à protéines, et on détecte la présence de l'antigène correspondant, spécifique de Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou d'un micro-organisme associé.

Une puce à protéines ci-dessus décrite peut être utilisée pour la détection de produits de gènes, pour établir un profil d'expression desdits gènes, en complément d'une puce à ADN selon l'invention.

Les puces à protéines selon l'invention sont également extrêmement utiles pour les expériences de protéomique, qui étudie les interactions entre les différentes protéines d'un micro-organisme donné. De façon simplifiée, on fixe des peptides représentatifs



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des différentes protéines d'un organisme sur un support. Puis, on met ledit support en contact avec des protéines marquées, et après une étape optionnelle de rinçage, on détecte des interactions entre lesdites protéines marquées et les peptides fixés sur la puce à protéines.

Ainsi, les puces à protéines comprenant une séquence polypeptidique selon l'invention ou un anticorps selon l'invention sont objet de l'invention, ainsi que les kits ou nécessaires les contenant.

La présente invention couvre également un procédé de détection et/ou d'identification de bactéries appartenant à l'espèce Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou à un micro-organisme associé dans un échantillon biologique, qui met en œuvre une séquence nucléotidique selon l'invention.

Il doit être entendu que le terme échantillon biologique concerne dans la présente invention les échantillons prélevés à partir d'un organisme vivant (en particulier sang, tissus, organes ou autres prélevés à partir d'un mammifère) ou un échantillon contenant du matériel biologique, c'est-à-dire de l'ADN ou de l'ARN. Un tel échantillon biologique comprend aussi les compositions alimentaires contenant des bactéries (par exemple les fromages, les produits laitiers), mais également des compositions alimentaires contenant des levures (bières, pains) ou autres. Le terme échantillon biologique concerne aussi les bactéries isolées à partir de ces prélèvements ou compositions alimentaires.

Le procédé de détection et/ou d'identification mettant en œuvre les séquences nucléotidiques selon l'invention peut être de diverse nature.

On préfère un procédé comportant les étapes suivantes :

- a) éventuellement, isolement de l'ADN à partir de l'échantillon biologique à analyser, ou obtention d'un ADNc à partir de l'ARN de l'échantillon biologique;
- b) amplification spécifique de l'ADN de bactéries appartenant à l'espèce Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou à un micro-organisme associé à l'aide d'au moins une amorce selon l'invention;
- c) mise en évidence des produits d'amplification.

Ce procédé est basé sur l'amplification spécifique de l'ADN, en particulier par une réaction d'amplification en chaîne.

On préfère également un procédé comprenant les étapes suivantes :

a) mise en contact d'une sonde nucléotidique selon l'invention avec un échantillon biologique, l'acide nucléique contenu dans l'échantillon biologique ayant, le cas

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échéant, préalablement été rendu accessible à l'hybridation, dans des conditions permettant l'hybridation de la sonde à l'acide nucléique d'une bactérie appartenant à l'espèce *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou à un microorganisme associé;

b) mise en évidence de l'hybride éventuellement formé entre la sonde nucléotidique et
 l'ADN de l'échantillon biologique.

Un tel procédé ne doit pas être limité à la détection de la présence de l'ADN contenu dans l'échantillon biologique à tester, il peut être également mis en œuvre pour détecter l'ARN contenu dans ledit échantillon. Ce procédé englobe en particulier les Southern et Northern blot.

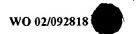
Un autre procédé préféré selon l'invention comprend les étapes suivantes :

- a) mise en contact d'une sonde nucléotidique immobilisée sur un support selon l'invention avec un échantillon biologique, l'acide nucléique de l'échantillon, ayant, le cas échéant, été préalablement rendu accessible à l'hybridation, dans des conditions permettant l'hybridation de la sonde à l'acide nucléique d'une bactérie appartenant à l'espèce Streptococcus agalactiae CIP 82.45 (ATCC 12403) ou à un micro-organisme associé;
- b) mise en contact de l'hybride formé entre la sonde nucléotidique immobilisée sur un support et l'acide nucléique contenu dans l'échantillon biologique, le cas échéant après élimination de l'ADN de l'échantillon biologique n'ayant pas hybridé avec la sonde, avec une sonde nucléotidique marquée selon l'invention;
- c) mise en évidence du nouvel hybride formé à l'étape b).

Ce procédé est avantageusement utilisé avec une puce à ADN selon l'invention, l'acide nucléique recherché s'hybridant avec une sonde présente à la surface de ladite puce, et étant détecté par l'utilisation d'une sonde marquée. Ce procédé est avantageusement mis en œuvre en combinant une étape préalable d'amplification de l'ADN ou de l'ADN complémentaire obtenu éventuellement par transcription inverse, à l'aide d'amorces selon l'invention.

Ainsi, la présente invention englobe également les kits ou nécessaires pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus* agalactiae CIP 82.45 (ATCC 12403) ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :

a) une sonde nucléotidique selon l'invention;



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- b) éventuellement, les réactifs nécessaires à la mise en œuvre d'une réaction d'hybridation;
- c) éventuellement, au moins une amorce selon l'invention ainsi que les réactifs nécessaires à une réaction d'amplification de l'ADN.

De même, la présente invention englobe également les kits ou nécessaires pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :

- a) une sonde nucléotidique, dite sonde de capture, selon l'invention ;
- b) une sonde oligonucléotidique, dite sonde de révélation, selon l'invention ;
  - c) éventuellement, au moins une amorce selon l'invention ainsi que les réactifs nécessaires à une réaction d'amplification de l'ADN.

Enfin, les kits ou nécessaires pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :

- a) au moins une amorce selon l'invention;
- b) éventuellement, les réactifs nécessaires pour effectuer une réaction d'amplification d'ADN;
- c) éventuellement, un composant permettant de vérifier la séquence du fragment amplifié, plus particulièrement une sonde oligonucléotidique selon l'invention, sont également objet de la présente invention.

De préférence, lesdites amorces et/ou sondes et/ou polypeptides et/ou anticorps selon la présente invention utilisés dans les procédés et/ou kits ou nécessaires selon la présente invention sont choisis parmi les amorces et/ou sondes et/ou polypeptides et/ou anticorps spécifiques de l'espèce *Streptococcus agalactiae* CIP 82.45 (ATCC 12403). De manière préférée, ces éléments sont choisis parmi les séquences nucléotidiques codant pour une protéine sécrétée, parmi les polypeptides sécrétés, ou parmi les anticorps dirigés contre des polypeptides sécrétés de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403).

La présente invention a également pour objet les souches de *Streptococcus* agalactiae CIP 82.45 (ATCC 12403) et/ou de micro-organismes associés contenant une ou plusieurs mutation(s) dans une séquence nucléotidique selon l'invention, en particulier une séquence ORF, ou leurs éléments régulateurs (en particulier promoteurs).

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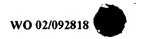
On préfère, selon la présente invention, les souches de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) présentant une ou plusieurs mutation(s) dans les séquences nucléotidiques codant pour des polypeptides impliqués dans la machinerie cellulaire, en particulier la sécrétion, le métabolisme intermédiaire central, le métabolisme énergétique, les processus de synthèse des acides aminés, de transcription et de traduction, de synthèse des polypeptides.

Lesdites mutations peuvent mener à une inactivation du gène, ou en particulier lorsqu'elles sont situées dans les éléments régulateurs dudit gène, à une surexpression de celui-ci.

L'invention concerne en outre l'utilisation d'une séquence nucléotidique selon l'invention, d'un polypeptide selon l'invention, d'un anticorps selon l'invention, d'une cellule selon l'invention, et/ou d'un animal transformé selon l'invention, pour la selection de composé organique ou inorganique capable de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, et/ou de modifier la réplication cellulaire de cellules eucaryotes ou procaryotes ou capables d'induire, d'inhiber ou d'aggraver une pathologies liées à une infection par *Streptococcus agalactiae* ou un de ses microorganismes associés.

L'invention comprend également une méthode de sélection de composés capables de se lier à un polypeptide ou un de ses fragments selon l'invention, capables de se lier à une séquence nucléotidique selon l'invention, ou capables de reconnaître un anticorps selon l'invention, et/ou capables de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, et/ou de modifier la croissance ou la réplication cellulaire de cellules eucaryotes ou procaryotes, ou capables d'induire, d'inhiber ou d'aggraver chez un organisme animal ou humain une pathologie liée à une infection par *Streptococcus*, par exemple par *Streptococcus agalactiae*, ou un de ses micro-organismes associés, caractérisée en ce qu'elle comprend les étapes suivantes :

- a) mise en contact dudit composé avec ledit polypeptide, ladite séquence nucléotidique, avec une cellule transformée selon l'invention et/ou administration dudit composé à un animal transformé selon l'invention;
- b) détermination de la capacité dudit composé à se lier avec ledit polypeptide ou ladite séquence nucléotidique, ou de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, ou de moduler la croissance ou la réplication cellulaire, ou d'induire, d'inhiber ou d'aggraver chez ledit animal transformé les pathologies liées



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à une infection par Streptococcus, par exemple Streptococcus agalactiae ou un de ses micro-organismes associés.

Les cellules et/ou les animaux transformés selon l'invention, pourront avantageusement servir de modèle et être utilisés dans des procédés pour étudier, identifier et/ou sélectionner des composés susceptibles d'être responsables de pathologies induites ou aggravées par *Streptococcus agalactiae*, ou susceptibles de prévenir et/ou de traiter ces pathologies. En particulier, les cellules hôtes transformées, notamment les bactéries de la famille des *Streptococcus* dont la transformation par un vecteur selon l'invention peut par exemple accroître ou inhiber son pouvoir infectieux, ou moduler les pathologies habituellement induites ou aggravées par l'infection, pourront être utilisées pour infecter des animaux dont on suivra l'apparition des pathologies. Ces animaux non transformés, infectés par exemple avec des bactéries *Streptococcus* transformées, pourront servir de modèle d'étude. De la même manière, les animaux transformés selon l'invention pourront être utilisés dans des procédés de sélection de composés susceptibles de prévenir et/ou de traiter les maladies dues à *Streptococcus*. Lesdits procédés utilisant lesdites cellules transformées et/ou animaux transformés, font partie de l'invention.

Les composés susceptibles d'être sélectionnés peuvent être des composés organiques tels que des polypeptides ou hydrates de carbone ou tous autres composés organiques ou inorganiques déjà connus, ou des composés organiques nouveaux élaborés à partir de techniques de modélisation moléculaire et obtenus par synthèse chimique ou biochimique, ces techniques étant connues de l'homme de l'art.

Lesdits composés sélectionnés pourront être utilisés pour moduler la croissance et/ou la réplication cellulaire de *Streptococcus agalactiae* ou tout autre micro-organisme associé et ainsi pour contrôler l'infection par ces micro-organismes. Lesdits composés selon l'invention pourront également être utilisés pour moduler la croissance et/ou la réplication cellulaire de toutes cellules eucaryotes ou procaryotes, notamment les cellules tumorales et les micro-organismes infectieux, pour lesquelles lesdits composés s'avéreront actifs, les méthodes permettant de déterminer lesdites modulations étant bien connues de l'homme de l'art.

On entend désigner par composé capable de moduler la croissance d'un microorganisme tout composé permettant d'intervenir, de modifier, de limiter et/ou de réduire le développement, la croissance, la vitesse de prolifération et/ou la viabilité dudit microorganisme.

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Cette modulation peut être réalisée par exemple par un agent capable de se lier à une protéine et ainsi d'inhiber ou de potentialiser son activité biologique, ou capable de se lier à une protéine membranaire de la surface extérieure d'un micro-organisme et de bloquer la pénétration dudit micro-organisme dans la cellule hôte ou de favoriser l'action du système immunitaire de l'organisme infecté dirigé à l'encontre dudit micro-organisme. Cette modulation peut être également réalisée par un agent capable de se lier à une séquence nucléotidique d'un ADN ou ARN d'un micro-organisme et de bloquer par exemple l'expression d'un polypeptide dont l'activité biologique ou structurelle est nécessaire à la croissance ou à la reproduction dudit micro-organisme.

On entend désigner par micro-organisme associé dans la présente invention, tout micro-organisme dont l'expression de gène peut être modulée, régulée, induite ou inhibée, ou dont la croissance ou la réplication cellulaire peut être également modulée par un composé de l'invention. On entend désigner également par micro-organisme associé dans la présente invention, tout micro-organisme comportant des séquences nucléotidiques ou des polypeptides selon l'invention. Ces micro-organismes peuvent dans certains cas comporter des polypeptides ou des séquences nucléotidiques identiques ou homologues à celles de l'invention et pourront également être détectés et/ou identifiés par les procédés ou kit de détection et/ou d'identification selon l'invention et également servir de cible pour les composés de l'invention. On entend aussi désigner par micro-organisme tout micro-organisme *Streptococcus agalactiae* de tout sérotype.

L'invention concerne les composés susceptibles d'être sélectionnés par une méthode de sélection selon l'invention.

L'invention concerne également une composition pharmaceutique comprenant un composé choisi parmi les composés suivants :

- a) une séquence nucléotidique selon l'invention;
- b) un polypeptide selon l'invention;
- c) un vecteur selon l'invention;
- d) un anticorps selon l'invention; et
- e) un composé susceptible d'être sélectionné par une méthode de sélection selon l'invention,

éventuellement en association avec un véhicule pharmaceutiquement acceptable.



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La présente invention concerne en outre une composition pharmaceutique selon l'invention pour la prévention et le traitement d'une infection par une bactérie appartenant à l'espèce Streptococcus agalactiae.

La présente invention concerne en outre une composition pharmaceutique selon l'invention, caractérisée en ce qu'elle comprend des anticorps dirigés contre les polypeptides spécifiques de *Streptococcus agalactiae*.

On entend désigner par quantité efficace, une quantité suffisante dudit composé ou anticorps, ou de polypeptide de l'invention, permettant de moduler la croissance de Streptococcus agalactiae ou d'un micro-organisme associé.

L'invention concerne aussi une composition pharmaceutique selon l'invention pour la prévention ou le traitement d'une infection par une bactérie appartenant au genre Streptococcus ou par un micro-organisme associé.

L'invention vise en outre une composition immunogène et/ou vaccinale, caractérisée en ce qu'elle comprend un ou plusieurs polypeptides selon l'invention et/ou un ou plusieurs polypeptides hybrides selon l'invention.

L'invention comprend aussi l'utilisation d'une cellule transformée selon l'invention, pour la préparation d'une composition vaccinale.

L'invention vise également une composition vaccinale, caractérisée en ce qu'elle contient une séquence nucléotidique selon l'invention, un vecteur selon l'invention et/ou une cellule transformée selon l'invention.

L'invention concerne en outre une composition immunogène capable d'induire une réponse immunitaire cellulaire ou humorale pour la prévention ou le traitement d'une infection par une bactérie appartenant à l'espèce *Streptococcus agalactiae*, caractérisée en ce qu'elle comprend une composition immunogène ou une composition vaccinale selon l'invention, en association avec un véhicule pharmaceutiquement acceptable et éventuellement un ou plusieurs adjuvants de l'immunité appropriés.

L'invention concerne également les compositions vaccinales selon l'invention, pour la prévention ou le traitement d'une infection par une bactérie appartenant au genre Streptococcus ou par un micro-organisme associé.

De manière préférée, les compositions immunogènes et/ou vaccinales selon l'invention destinées à la prévention et/ou au traitement d'infection par *Streptococcus* ou par un micro-organisme associé seront choisies parmi les compositions immunogènes et/ou vaccinales comprenant un polypeptide ou un de ses fragments correspondant à une protéine, ou un de ses fragments, de l'enveloppe cellulaire de *Streptococcus*. Les

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compositions vaccinales comprenant des séquences nucléotidiques comprendront de préférence également des séquences nucléotidiques codant pour un polypeptide ou un de ses fragments correspondant à une protéine, ou un de ses fragments, de l'enveloppe cellulaire de *Streptococcus*.

Les polypeptides de l'invention ou leurs fragments entrant dans les compositions immunogènes selon l'invention peuvent être sélectionnés par des techniques connues de l'homme de l'art comme par exemple sur la capacité desdits polypeptides à stimuler les cellules T, qui se traduit par exemple par leur prolifération ou la sécrétion d'interleukines, ou qui aboutit à la production d'anticorps dirigés contre lesdits polypeptides.

Chez la souris, chez laquelle une dose pondérale de la composition vaccinale comparable à la dose utilisée chez l'homme est administrée, la réaction anticorps est testée par prélèvement du sérum suivi d'une étude de la formation d'un complexe entre les anticorps présents dans le sérum et l'antigène de la composition vaccinale, selon les techniques usuelles.

Selon l'invention, lesdites compositions vaccinales seront de préférence en association avec un véhicule pharmaceutiquement acceptable et, le cas échéant, avec un ou plusieurs adjuvants de l'immunité appropriés.

Aujourd'hui, divers types de vaccins sont disponibles pour protéger l'homme contre des maladies infectieuses : micro-organismes vivants atténués (M. bovis - BCG pour la tuberculose), micro-organismes inactivés (virus de la grippe), des extraits acellulaires (Bordetella pertussis pour la coqueluche), protéines recombinées (antigène de surface du virus de l'hépatite B), des polyosides (pneumocoques). Des vaccins préparés à partir de peptides de synthèse ou de micro-organismes génétiquement modifiés exprimant des antigènes hétérologues sont en cours d'expérimentation. Plus récemment encore, des ADNs plasmidiques recombinés portant des gènes codant pour des antigènes protecteurs ont été proposés comme stratégie vaccinale alternative. Ce type de vaccination est réalisé avec un plasmide particulier dérivant d'un plasmide de E. coli qui ne se réplique pas in vivo et qui code uniquement pour la protéine vaccinante. Des animaux ont été immunisés en injectant simplement l'ADN plasmidique nu dans le muscle. Cette technique conduit à l'expression de la protéine vaccinale in situ et à une réponse immunitaire de type cellulaire (CTL) et de type humoral (anticorps). Cette double induction de la réponse immunitaire est l'un des principaux avantages de la technique de vaccination avec de l'ADN nu.

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Les compositions vaccinales comprenant des séquences nucléotidiques ou des vecteurs dans lesquels sont insérées lesdites séquences, sont notamment décrites dans la demande internationale N° WO 90/11092 et également dans la demande internationale N° WO 95/11307.

La séquence nucléotidique constitutive de la composition vaccinale selon l'invention peut être injectée à l'hôte après avoir été couplée à des composés qui favorisent la pénétration de ce polynucléotide à l'intérieur de la cellule ou son transport jusqu'au noyau cellulaire. Les conjugués résultants peuvent être encapsulés dans des microparticules polymères, comme décrit dans la demande internationale N° WO 94/27238 (Medisorb Technologies International).

Selon un autre mode de réalisation de la composition vaccinale selon l'invention, la séquence nucléotidique, de préférence un ADN, est complexée avec du DEAE-dextran, avec des protéines nucléaires, avec des lipides ou encapsulée dans des liposomes ou encore introduite sous la forme d'un gel facilitant sa transfection dans les cellules. Le polynucléotide ou le vecteur selon l'invention peut aussi être en suspension dans une solution tampon ou être associé à des liposomes.

Avantageusement, un tel vaccin sera préparé conformément à la technique décrite par Tacson et al. ou Huygen et al. en 1996 ou encore conformément à la technique décrite par Davis et al. dans la demande internationale N° WO 95/11307.

Un tel vaccin peut être également préparé sous la forme d'une composition contenant un vecteur selon l'invention, placée sous le contrôle d'éléments de régulation permettant son expression chez l'homme ou l'animal. On pourra par exemple utiliser, en tant que vecteur d'expression *in vivo* de l'antigène polypeptidique d'intérêt, le plasmide pcDNA3 ou le plasmide pcDNA1/neo, tous les deux commercialisés par Invitrogen (R & D Systems, Abingdon, Royaume-Uni). Un tel vaccin comprendra avantageusement, outre le vecteur recombinant, une solution saline, par exemple une solution de chlorure de sodium.

On entend désigner par véhicule pharmaceutiquement acceptable, un composé ou une combinaison de composés entrant dans une composition pharmaceutique ou vaccinale ne provoquant pas de réactions secondaires et qui permet par exemple la facilitation de l'administration du composé actif, l'augmentation de sa durée de vie et/ou de son efficacité dans l'organisme, l'augmentation de sa solubilité en solution ou encore l'amélioration de sa conservation. Ces véhicules pharmaceutiquement acceptables sont

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bien connus et seront adaptés par l'homme de l'art en fonction de la nature et du mode d'administration du composé actif choisi.

En ce qui concerne les formulations vaccinales, celles-ci peuvent comprendre des adjuvants de l'immunité appropriés qui sont connus de l'homme de l'art, comme par exemple l'hydroxyde d'aluminium, un représentant de la famille des muramyl peptides comme un des dérivés peptidiques du N-acétyl-muramyl, un lysat bactérien, ou encore l'adjuvant incomplet de Freund.

De préférence, ces composés seront administrés par voie systémique, en particulier par voie intraveineuse, par voie intramusculaire, intradermique ou souscutanée, ou par voie orale. De manière plus préférée, la composition vaccinale comprenant des polypeptides selon l'invention, sera administrée à plusieurs reprises, de manière étalée dans le temps, par voie intradermique ou sous-cutanée.

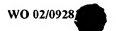
Leurs modes d'administration, posologies et formes galéniques optimaux peuvent être déterminés selon les critères généralement pris en compte dans l'établissement d'un traitement adapté à un patient comme par exemple l'âge ou le poids corporel du patient, la gravité de son état général, la tolérance au traitement et les effets secondaires constatés.

Enfin, l'invention comprend l'utilisation d'une composition selon l'invention, pour le traitement ou la prévention de maladies induites ou aggravées par la présence de Streptococcus.

Par ailleurs, la présente invention a également pour objet une banque d'ADN génomique d'une bactérie du genre *Streptococcus*, de manière préférée, *Streptococcus* agalactiae, de manière préférée la CIP 82.45 (ATCC 12403).

La banque d'ADN génomique décrite dans la présente invention, en particulier la banque déposée à la CNCM le 28 décembre 2000 sous le numéro d'ordre N° I-2610, recouvre en effet *Streptococcus agalactiae* CIP 82.45 (ATCC 12403). Toutefois, si certaines régions n'ont pas pu être clonées dans ladite banque, en raison de problèmes de létalités chez *Escherichia coli*, ces régions peuvent facilement être amplifiées et identifiées par l'homme du métier, en utilisant des oligonucléotides spécifiques des séquences des extrémités des différents clones qui forment les contigs.

La présente invention concerne également les méthodes pour l'isolement d'un polynucléotide d'intérêt présent chez une souche de *Streptococcus agalactiae* et absente chez une autre souche, qui utilise au moins une banque d'ADN basée par exemple sur un plasmide pSYX34 contenant un fragment du génome de *Streptococcus agalactiae*.



La méthode selon l'invention pour l'isolement d'un polynucléotide d'intérêt peut comprendre les étapes suivantes :

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 a) isoler au moins un polynucléotide contenu dans un clone de la banque d'ADN d'origine de Streptococcus agalactiae,

## 5 b) isoler:

- au moins un polynucléotide génomique ou ADNc d'une bactérie Streptococcus, ladite bactérie Streptococcus appartenant à une souche différente de la souche utilisée pour la construction de la banque d'ADN de l'étape a) ou, de façon alternative,
- au moins un polynucléotide contenu dans un clone d'une banque d'ADN préparé à partir du génome d'une Streptococcus qui est différente de la souche Streptococcus agalactiae utilisée pour la construction de la banque d'ADN de l'étape a);
  - c) hybrider le polynucléotide de l'étape a) au polynucléotide de l'étape b);
  - d) sélectionner les polynucléotides de l'étape a) qui n'ont pas formé de complexe d'hybridation avec les polynucléotides de l'étape b);
    - e) caractériser le polynucléotide sélectionné.

On peut préparer le polynucléotide de l'étape a) par la digestion d'au moins un clone recombinant avec une enzyme de restriction appropriée, et de façon optionnelle, l'amplification de l'insert polynucléotide qui en résulte.

Ainsi, la méthode de l'invention permet à l'homme du métier d'effectuer des études génomiques comparatives entre les différentes souches ou espèces du genre *Streptococcus*, par exemple entre les souches pathogéniques et leurs équivalents non pathogènes.

En particulier, il est possible d'étudier et de déterminer les régions de polymorphisme entre lesdites souches.

L'invention comprend aussi un procédé d'identification de séquence spécifique de *Streptococcus agalactiae*, caractérisé par l'alignement de séquences nucléotidiques de *Streptococcus agalactiae* selon l'invention et le traitement de données obtenues par cet alignement pour isoler les séquences spécifiques.

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La présente invention concerne également l'utilisation des séquences nucléiques ou des polypeptides selon la présente invention :

- pour la sécrétion de protéines,
- comme facteurs de virulence,

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- pour le contrôle par l'intermédiaire du quorum-sensing,
- pour l'identification de cibles pour les maladies humaines dont *Streptococcus* agalactiae est un modèle, et
- pour l'identification de cibles contre les bactéries Gram positives pathogènes par la méthode de génomique soustractive (comme par exemple par comparaison avec des bactéries Gram positives non pathogènes).

#### **EXEMPLES**

# Exemple 1: Matériels et méthode

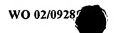
La stratégie de séquençage du génome de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) repose sur un séquençage alléatoire (shot-gun). La première étape de ce travail consiste à cloner l'ADN génomique de la bactérie *Streptococcus agalactiae* dans differents vecteurs (plasmides et BAC).

Matériels et méthodes.

- 1. Construction des banques :
  - a/ Banque de petits fragments dans le vecteur pcDNA2.1

L'ADN chromosomique de la souche de Streptococcus agalactiae CIP 82.45 (ATCC 12403) a été préparé par une méthode classique incluant un traitement à la protéinase K et une extraction au phénol (9). Environ 10 μg d'ADN ont été cassés par nébulisation (1 minute sous une pression de 1 bar) (4). Les extrémités des fragments d'ADN ont été rendues franches en faisant agir la DNA-polymérase du bactériophage T4 pendant 15 minutes à 37°C en présence des 4 nucléotides tri-phosphate. L'enzyme a été inactivée par une incubation de 15 mn à 75°C. Des adaptateurs (invitrogen Cat. Nº 408-18) ont ensuite été ligaturés à ces extrémités. Après ligature, les fragments d'ADN chromosomiques ayant une taille entre 1 000 et 3 000 paires de bases ont été purifiés après électrophorèse sur gel d'agarose. Le vecteur utilisé pour la construction de la banque, pcDNA2.1 (Invitrogen), a été digéré par l'enzyme BstX1 et purifié par geneclean (BIO-101) après électrophorèse sur gel d'agarose. L'ADN chromosomique et le vecteur purifié ont été ligaturés par action de la ligase du bactériophage T4. Le mélange de ligation a été introduit par transformation dans la souche d'Escherichia coli XL2-blue (Stratagene). Environ 4000 colonies sont obtenues par µl du mélange de ligation.

b/ Construction d'une banque de fragments de taille moyenne (banque déposée à la CNCM sous le numéro I-2610)



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Afin de limiter la fréquence des événements de co-ligation et de clones sans insert, la méthode du remplissage partiel du site de coupure a été utilisée (7).

L'ADN chromosomique de la souche de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) a été digéré de manière partielle par l'enzyme de restriction Sau3A en utilisant une gamme de dilution de l'enzyme. La fraction présentant la gamme de tailles souhaitée (entre 2 et 12 kbases) a été précipitée et les extrémités ont été remplies partiellement par le fragment de Klenow de la DNA polymérase d'*E. coli* en présence de dTTP et dCTP. Après purification sur gel d'agarose les fragments d'ADN chromosomique ayant une taille comprise entre 3 et 6 kbases ont été ligués au vecteur pSYX34 (7) digéré par l'enzyme SalI remplie partiellement par l'enzyme de Klenow en présence de dATP et dGTP. Le mélange de ligation a été introduit par transformation dans la souche d'*Escherichia coli* XL10-kan (Stratagene) et étalement sur milieu LB contenant du chloramphénicol à une concentration de 20 mg/l. Environ 500 colonies sont obtenues par μl du mélange de ligation.

## 2. Préparation des plasmides et séquençage

Les plasmides ont été préparés par une méthode semi-automatique de préparation développée au laboratoire GMP basé sur la méthode de lyse alcaline (2). Les inserts chromosomiques ont été séquencés à partir de leurs deux extrémités en utilisant les primer T7 et universel en suivant les recommandations du fournisseur (PE-biosystems). Les séquences ont été déterminées en utilisant un séquenceur automatique de type 3700 (PE-Biosystem).

#### 3. Assemblage des séquences

Les séquences ont été assemblées en utilisant l'ensemble de logiciel développé à l'Université de Washington, Phred, Phrap et Consed (5, 8). La finition de la séquence a été réalisée en utilisant l'ensemble de logiciel GMPTB (7). L'étape de finition correspond au reséquençage des régions où la séquence est peu sûre et le séquençage des régions situées entre les contigs. Elle a été réalisée en séquençant des produits de PCR correspondant à ces régions identifiées par un expert en utilisant les logiciels Consed (8) et GMPTB (7). Les séquences des oligonucléotides ont été définies en utilisant les logiciels Consed et Primo (8, 10).

## 4. Annotation des séquences

L'identification des phases codantes (CDS) a été réalisée en utilisant l'ensemble de logiciel GMPTB (L. Frangeul et al. non publié). Ce programme combine les résultats de différentes méthodes : (i) l'identification de phases ouvertes de lecture et

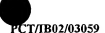
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leur tri en fonction de leur taille, (ii) l'analyse de la probabilité d'être codante en utilisant le logiciel Genemark (11), (iii) l'identification d'un début de traduction (codon d'initiation et séquence de fixation du ribosome), (iv) similarité de la séquence protéique déduite avec les séquences protéiques contenues dans les banques de séquence en utilisant le logiciel BLASTP.

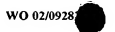
Les fonctions des protéines codées par les phases codantes identifiées indiquées au tableau 1 ont été prédites par l'analyse des résultats de recherche de similarités dans les banques en utilisant le logiciel BLASTP (1).

Exemple 2: Description scientifique de la banque de BAC de Streptococcus agalactiae CIP 82.45 (ATCC 12403) déposée à la CNCM le 28 décembre 2000 sous le numéro I-2610.

Collection de clones d'Escherichia coli DH 10B™ (Calvin et al., J. Bacteriol. 170, 2796, 1988) contenant des fragments d'ADN génomique de la bactérie Streptococcus agalactiae souche NEM 316,CIP 82.45 (ATCC 12403), clonés dans le vecteur pSYX34 (Xu et al., Biotechniques, 17:57, 1990). Le vecteur a été digéré SalI et partiellement rempli à l'aide de l'ADN polymérase afin de produire des extrémités 5'-TC. Des fragments génomiques aléatoires de 3-6 kb provenant de Streptococcus agalactiae (souche NEM 316, CIP 82-45, ATCC 12403) ont été digérés partiellement par Sau3A et partiellement remplis afin de produire des extrémités 5'-GA. Après ligature in vitro et transformation, des clones résistant au chloromphénicol ont été sélectionnés. Environ 5000 clones ont été assemblés, suspendus dans 15 ml de milieu L et congelés.

Exemple 3 : Les protéines de surface de Streptococcus agalactiae NEM316

Les protéines de surface des bactéries pathogènes, et plus particulièrement les protéines dites du type LPXTG (Navarre et Schneewind, Microbial. Mol. Biol. Rev. 63 174-229), jouent un rôle crucial lors du processus infectieux en permettant notamment des interactions entre le microorganisme et les cellules de l'hôte et/ou l'échappement au système immunitaire. Les inventeurs ont don focalisé leur étude sur ce type de protéines qui présentent la particularité d'être liées covalemment au peptidoglycane via le motif d'ancrage carboxylique LPXTG. Cette réaction est catalysée par une enzyme bifonctionnelle (endopeptidase-transpeptidase) appelée sortase. L'étude du rôle de ces protéines dans la virulence de S. agalactiae a été réalisée par 2 approches



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complémentaires (construction d'un mutant sortase-déficient, inactivation des gènes codant pour des protéines de type LPXTG).

- Le gène srtA de S. agalactiae NEM316 (IPF N°1268).

Le mutant MEM1979, déposé le 24 avril 2002 à la CNCM sous le numéro I-2861, est une souche mutante dérivée de NEM316 (CIP 82.45, ATCC 12403) dans laquelle l'IPF 1268 a été inactivé.

L'analyse du génome de NEM316 nous a permis de caractériser un gène srtA présentant homologie respectivement 55 % et 30 % d'identité avec les sortases de Streptococcus gordonii et de Staphylococcus aureus. Ce gène a été inactivé par insertion-inactivation et nous avons montré que le mutant ainsi construit n'adhérait plus à des cellules épithéliales humaines pulmonaires (A549) et utérines (HeLa). Ces résultats suggèrent que les protéines du type LPXTG de S. agalactiae jouent un rôle dans la virulence de cette bactérie, en permettant notamment son adhésion avec les cellules de l'hôte.

- Recensement des protéines du type LPXTG de S. agalactiae NEM316.

Une analyse in silico du génome de NEM316 a révélé la présence de 30 protéines de surface putatives possédant le motif d'ancrage de type LPXTG (Tableau 6). Nous avons étudié par amplification PCR, en utilisant des amorces spécifiques, la distribution des gènes codant pour 21 protéines du type LPXTG dans une collection de 99 souches non-redondantes de S. agalactiae responsables d'infections non-invasives (70 souches provenant de portage ou d'infection urinaire) et invasives (29 souches provenant d'hémoculture ou de méningite). Cette étude a montré que 6 de ces gènes (IPN N° 1503, 678, 2192, 1861, 584, 280) étaient présents dans toutes les souches de notre échantillonnage (Tableau 7). Deux de ces six gènes (IPF N°678 et 1503) ont été inactivés par insertion-inactivation et les mutants correspondants NEM2056 et NEM2057 présentaient une adhérence diminuée avec les cellules épithéliales A549 (Tableau 5).

Le mutant NEM2056, déposé le 24 avril 2002 à la CNCM sous le numéro I-2862, est une souche mutante dérivée de NEM316 (CIP 82.45, ATCC 12403) dans laquelle l'IPF 678 a été inactivé.

Le mutant NEM2057, déposé le 24 avril 2002 à la CNCM sous le numéro I-2863, est une souche mutante dérivée de NEM316 (CIP 82.45, ATCC 12403) dans laquelle l'IPF 1503 a été inactivé.



- Cibles vaccinales.

La présence des gènes IPF N°1503, 678, 2192, 1861, 584, 280 chez toutes les souches de S. agalactiae testées font des protéines correspondantes des cibles vaccinales de choix pour le développement d'un vaccin anti-S. agalactiae.

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<u>TABLEAU 1</u>: Liste des phases codantes annotées identifiées par l'analyse des séquences des contigs

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SEQ ID No.	Prot No.	Localisation et sens sur contig	Résultats redondantes & Homologie	Résultats Blastp sur des banques de protéines non ndantes mologie
SeqID 140	SA-1.2	Contig137 (42738-43340 p)	74	Identities = 129/202 (63%), Positives = 156/202 (76%), Gaps = 8/202 (3%) ref[NP_072029.1] traG-related protein [Enterococcus faecalis] gblAAG40447.1  (AE002565) traG-related protein [Enterococcus faecalis]
SeqID 141	SA-10.1	Contig137 (33454-33681 m)	No Hits found	
SeqID 142	SA-1000.1	Contig127 (13387-14454 p)	75	Identities = 220/354 (62%), Positives = 274/354 (77%), Gaps = 1/354 (0%) splQ48677 PEPA_LACLC GLUTAMYL-AMINOPEPTIDASE emb CAA56994.1  (X81089) glutamylaminopeptidase [Lactococcus lactis] Length = 355
SeqID 143	SA-1001.1	Contig127 (12912-13202 p)	No Hits found	
SeqID 144	SA-1002.1	Contig127 (12518-12802 m)	No Hits found	
SeqID 145	SA-1003.1	Contig127 (12198-12407 m)	65	Identities = 49/100 (49%), Positives = 71/100 (71%), Gaps = 4/100 (4%) dbj BAB06972.1  (AP001518) thioredoxin H1 [Bacillus halodurans]
SeqID 146	SA-1004.1	Contig127 (11539-12165 m)	09	Identities = 79/196 (40%), Positives = 126/196 (63%), Gaps = 1/196 (0%) pir  A69999 phenylalanyl-tRNA synthetase (beta subunit) homolog ytpR - Bacillus subtilis gb AAC00291.1  (AF008220) YtpR [Bacillus subtilis] emb CAB14960.1  (299119) similar to phenylalanyl-tRNA synthetase (beta subunit) [Bacillus subtilis] Length = 201
SeqID 147	SA-1006.1	Contig127 (10769-11485 p)	47	Identities = 69/243 (28%), Positives = 117/243 (47%), Gaps = 15/243 (6%) sp Q02148 YHI6_LACLA HYPOTHETICAL 30.7 KDA PROTEIN IN HISD-HISB INTERGENIC REGION (ORF6) pir  F45734 orf6 3 to hisD - Lactococcus lactis subsp. lactis gb AAB81904.1  (U92974) unknown {Lactococcus lactis} Length = 263



/141 (65%), Gaps = 45) putative single lactis bacteriophage 41	= 231/279 (82%) porter ATP-binding Length = 296	/179 (50%), Gaps = 3) putative hydrolase Length = 238	ives = 371/585 (63%), Gaps = sonent sensor histidine kinase subtilis emb CAA99610.1  kinase [Bacillus subtilis] -component sensor histidine Length = 593	77244 (64%), Gaps = sponse regulator lytT1 (Z75208) autolysin 2AB14852.1 (Z99118) Bacillus subtilis]		//110 (74%), Gaps = netical protein ysbA - 8) hypothetical protein 299118) similar to [Bacillus subtilis]
Identities = 65/141 (46%), Positives = 93/141 (65%), Gaps = 10/141 (7%) gb AAF74079.1  (AF212845) putative single stranded binding protein [Lactococcus lactis bacteriopha ul36] Length = 141	Identities = 201/279 (72%), Positives = 231/279 (82%) gbJAAC61484.1  (AF082738) ABC transporter ATP-binding protein [Streptococcus pyogenes] Length = 296	Identities = 63/179 (35%); Positives = 91/179 (50%), Gaps = 2/179 (1%) emb[CAC13072.1] (AL445503) putative hydrolase [Streptomyces coellcolor A3(2)]	Identities = 254/585 (43%), Positives = 371/585 (63%), Gaps = 9/585 (1%) pir[ A69655 two-component sensor histidine kinase lytS-involved - Bacillus subtilis emb CAA99610.1  (Z75208) autolysin sensor kinase [Bacillus subtilis] emb CAB14853.1  (Z99118) two-component sensor histidine kinase [Bacillus subtilis]	Identities = 105/244 (43%), Positives = 157/244 (64%), Gaps = 6/244 (2%) pir  B69655 two-component response regulator lytT-involved - Bacillus subtilis emb CAA99611.1  (Z75208) autolysin response regulator [Bacillus subtilis] emb CAB14852.1  (Z99118) two-component response regulator [Bacillus subtilis] Length = 241		Identities = 50/110 (45%), Positives = 82/110 (74%), Gaps = 2/110 (1%) pir  C69983 conserved hypothetical protein ysbA - Bacillus subtilis emb CAA99612.1  (Z75208) hypothetical protein [Bacillus subtilis] emb CAB14851.1  (Z99118) similar to hypothetical proteins from B. subtilis [Bacillus subtilis] Length = 146
69	82	33	09		No Hits found	51
Contig127 (10293-10688 m)	Contig131 (21108-21947 p)	Contig127 (9526-10170 m)	Contig127 (7754-9499 m)	Contig127 (7033-7779 m)	Contig127 (6936-7100 p)	Contig127 (6408-6863 m)
SA-1007.1	SA-101.1	SA-1010.1	SA-1012.1	SA-1013.1	SA-1014.1	SA-1016.1
SeqID 148	SeqID 149	SeqID 150	SeqID 151	SeqID 152	SeqID 153	SeqID 154

SeqID 155	SA-1017.1	Contig127 (5678-6406 m)	2	Identities = 120/240 (50%). Positives = 159/240 (66%), Gaps = 10/240 (4%) pir[:D69983 conserved hypothetical protein ysbB - Bacillus subtitis emb[CAA99613 1] (275208) hypothetical protein [Bacillus subtilis] emb[CAB14850.1] (299118) similar to hypothetical proteins [Bacillus subtilis]
SeqID 156	SA-1018.1	Contig127 (3807-5435 m)	06	Identities = 498/542 (91%), Positives = 518/542 (94%) gb AAC67217.1  (U78968) surface lipoprotein DppA [Streptococcus pyogenes] Length = 542
SeqID 157	SA-1019.1	Contig127 (2717-3694 m)	96	Identities = 302/325 (92%); Positives = 317/325 (96%) gb AAC67218.1  (U78968) transmembrane transport protein DppB [Streptococcus pyogenes] Length = 325
SeqID 158	SA-102.1	Contig131 (20566-21108 p)	69	Identities = 119/165 (72%), Positives = 134/165 (81%), Gaps = 5/165 (3%) gb AAC61483.1  (AF082738) phosphotidylglycerophosphate synthase [Streptococcus pyogenes] Length = 165
SeqID 159	SA-1020.1	Contig127 (1899-2720 m)	85	Identities = 255/274 (93%), Positives = 263/274 (95%), Gaps = 1/274 (0%) gb[AAC67219.1] (U78968) transmembrane transport protein DppC [Streptococcus pyogenes] Length = 274
SeqID.160	SA-1021.1	Contig127 (1084-1887 m)	95	Identities = 254/267 (95%), Positives = 262/267 (97%) gb AAC67220.1  (U78968) ATPase protein DppD [Streptococcus pyogenes] Length = 267
SeqID 161	SA-1022.1	Contig127 (474-1100 m)	91	Identities = 185/205 (90%), Positives = 195/205 (94%) gb AAC67221.1  (U78968) ATPase protein DppE [Streptococcus pyogenes] Length = 208



	Contig127 (1-192 m)	62	Identities = 35/58 (60%), Positives = 44/58 (75%) splP36672 PTTB_ECOL! PTS SYSTEM, TREHALOSE- SPECIFIC IIBC COMPONENT (EIIBC-TRE) (TREHALOSE ENZYME II, BC COMPONENT) (PHOSPHOTRANSFERASE phosphotransferase system trehalose permease - Escherichia coli (strain K-12) gblAAC77197.11 (AE000495) PTS system enzyme II, trehalose specific [Escherichia Length = 473
Contig113 (	(19147-19281 p)	No Hits found	
_	18773-19144 p)	53	Identities = 37/124 (29%), Positives = 67/124 (53%), Gaps = 2/124 (1%) gb AAC98903.1   (AF023181) DivIC homolog [Listeria monocytogenes] Length = 128
Contig113 (18	(18514-18786 p)	75	Identities = 49/84 (58%), Positives = 70/84 (83%) splP37557 YABO_BACSU HYPOTHETICAL 9.7 KDA PROTEIN IN MFD-DIVIC INTERGENIC REGION pir  S66089 conserved hypothetical protein yabO - Bacillus subtilis dbj BAA05294.1  (D26185) unknown [Bacillus subtilis] emb CAB11835.1  (Z99104) similar to hypothetical proteins [Bacillus subtilis] Length =
Contig113 (14	(14726-18223 p)	62	Identities = 523/1051 (49%), Positives = 733/1051 (68%), Gaps = 20/1051 (1%) gb AAD03810.1  (AF054624) transcription-repair coupling factor [Lactobacillus sakei] Length = 1045
Contig113 (1	(14154-14729 p)	9	Identities = 89/187 (47%), Positives = 127/187 (67%), Gaps = 2/187 (1%) dbjjBAB03787.1  (AP001507) stage V sporulation protein C (peptidyl-tRNA hydrolase) [Bacillus halodurans] Length = 185
Contig131 (1	(19160-20443 p)	. 62	dentities = 207/424 (48%), Positives = 277/424 (64%), Gaps = 3/424 (0%) gb AAC61481.1  (AF082738) unknown [Streptococcus pyogenes]

SeqID 169	SA-1030.1	Contig113 (12955-14070 p)	84	Identities = 261/371 (70%), Positives = 313/371 (84%), Gaps = 5/371 (1%) dbj BAB07770.1  (AP001520) GTP-binding protein [Bacillus halodurans] Length = 366
SeqID 170	SA-1031.1	Contig113 (12592-12795 p)	No Hits found	
SeqID 171	SA-1032.1	Contig113 (12193-12390 p)	82	Identities = 46/63 (73%), Positives = 57/63 (90%)   gb AAC45338.1  (AF000658) ORFX [Streptococcus pneumoniae]   Length = 64
SeqID 172	SA-1033.1	Contig113 (11290-12183 p)	43	Identities = 83/298 (27%), Positives = 138/298 (45%), Gaps = 35/298 (11%) pir  H6995 hypothetical protein ytlR - Bacillus subtilis gb AAC00282.1  (AF008220) YtlR [Bacillus subtilis] emb CAB14972.1  (Z99119) ytlR [Bacillus subtilis]
SeqID 173	SA-1034.1	Contig113 (10096-11232 p)	80	Identities = 278/378 (73%), Positives = 324/378 (85%) splO06672jDP3B_STRPN DNA POLYMERASE III, BETA CHAIN gb AAC45337.1  (AF000658) beta subunit of DNA polymerase III [Streptococcus pneumoniae] Length = 378
SeqID 174	SA-1035.1	Contig113 (8580-9941 p)	82	Identities = 353/455 (77%), Positives = 401/455 (87%), Gaps = 6/455 (1%) gb AAF71535.1 AF255728_1 (AF255728) chromosomal initiator protein DnaA [Streptococcus pyogenes] Length = 451
SeqID 175	SA-1036.1	Contig113 (7608-8381 p)	7.1	Identities = 138/257 (53%), Positives = 188/257 (72%), Gaps = 5/257 (1%) gb AAC45335.1  (AF000658) SPSpoJ [Streptococcus pneumoniae] Length = 252
SeqID 176	SA-1037.1	Contig113 (6275-7510 p)	0.2	Identities = 222/396 (56%), Positives = 290/396 (73%), Gaps = 27/396 (6%) gb AAC45334.1   (AF000658) putative serine protease [Streptococcus pneumoniae] Length = 397
SeqID 177	SA-1038.1	Contig113 (5601-6080 m)	. 08	Identities = 111/159 (69%), Positives = 136/159 (84%) gb AAC44894.1  (U76218) unknown [Streptococcus pneumoniae] gb AAC45340.1  (AF000658) ORF1 [Streptococcus pneumoniae] Length = 159



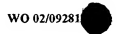
Contig113
Contig131 (17914-19158 p)
Contig113 (701-2320 p)
Contig113 (111-635 p)
Contig130 (3069-3740 p)
Contig130 (3867-5126 p)
Contig130 (5210-5782 p)
Contig130 (5763-7058 p)

SeqID 186	SA-1048.1	Contig130 (7081-7941 p)	44	Identities = 101/207 (48%), Positives = 128/207 (61%), Gaps = 31/207 (14%) gb AAC35914.1  (AF071085) methionine aminopeptidase A [Enterococcus faecalis] Length = 178
SeqiD 187	SA-1049.1	Contig130 (7943-8863 p)	49	Identities = 83/276 (30%), Positives = 157/276 (56%), Gaps = 3/276 (1%) gb AAC35915.1  (AF071085) Orfde2 [Enterococcus faecalis]
SeqID 188	SA-105.1	Contig131 (17322-17675 m)	61	Identities = 56/116 (48%), Positives = 74/116 (63%), Gaps = 3/116 (2%) pir  JC4754 hypothetical 13.6k protein - Lactococcus lactis emb CAA61548.1  (X89367) orf121 [Lactococcus lactis]
SeqID 189	SA-1050.1	Contig130 (8880-9314 m)	90	Identities = 52/161 (32%), Positives = 78/161 (48%), Gaps = 26/161 (16%) gb AAD54224.1 AF143443_3 (AF143443) MesH [Leuconostoc mesenteroides]
SeqID 190	SA-1051.1	Contig130 (9517-10026 p)	47	Identities = 56/158 (35%), Positives = 91/158 (57%), Gaps = 8/158 (5%) emb CAA68244.1  (X99978) citrulline cluster-linked gene [Lactobacillus plantarum] Length = 168
SeqID 191	SA-1053.1	Contig130 (10166-12124 p)	70	Identities = 345/657 (52%), Positives = 462/657 (69%), Gaps = 8/657 (1%) spl031498jDNLJ_BACSU DNA LIGASE (POLYDEOXYRIBONUCLEOTIDE SYNTHASE [NAD+]) pir[ F69794 DNA ligase (NAD+) (EC 6.5.1.2) - Bacillus subtilis emb CAB12482.1  (Z99107) similar to DNA ligase [Bacillus subtilis] Length = 668
SeqID 192	SA-1054.1	Contig130 (12136-13155 p)	54	Identities = 151/293 (51%), Positives = 193/293 (65%), Gaps = 5/293 (1%) pir  F69795 conserved hypothetical protein yerQ - Bacillus subtilis emb CAB12492.1  (299107) similar to hypothetical proteins [Bacillus subtilis] Length = 303
SeqID 193	SA-1055.1	Contig130 (13159-15459 p)	47	Identities = 227/669 (33%), Positives = 363/669 (53%), Gaps = 66/669 (9%) prf[1808262A thermostable pullulanase [Bacillus stearothermophilus] Length = 658

Identities = 213/412 (51%), Positives = 292/412 (70%), Gaps = 5/412 (1%) splP42086JPBUX_BACSU XANTHINE PERMEASE pir][S51310 xanthine transport protein pbuX - Bacillus subtilis emb]CAA58759.1] (X83878) xanthine permease [Bacillus subtilis] gb]AAA96612.1] (L77246) transport protein [Bacillus subtilis] emb]CAB14123.1] (Z99115) xanthine permease [Bacillus subtilis] Length = 438	Identities = 82/253 (32%), Positives = 119/253 (46%), Gaps = 10/253 (3%) splP39608 YWCJ_BACSU HYPOTHETICAL 28.4 KD PROTEIN IN SACT-SACP INTERGENIC REGION pir  S39703 nitrite transport protein homolog ywcJ - Bacillus subtilis emb CAA51604.1  (X73124) ipa-48r [Bacillus subtilis] emb CAB15832.1  (Z99123) alternate gene name: ipa-48r-similar to nitrite transporter [Bacillus subtilis]	Identities = 80/162 (49%), Positives = 112/162 (68%)   gb AAG18632.1  (AY007504) unknown [Streptococcus mitis]   Length = 173	Identities = 93/160 (58%), Positives = 120/160 (74%), Gaps = 1/160 (0%) gb AAG18632.1  (AY007504) unknown [Streptococcus mitis] Length = 173	Identities = 103/317 (32%), Positives = 165/317 (51%), Gaps = 15/317 (4%) spl083774 APBE_TREPA THIAMINE BIOSYNTHESIS LIPOPROTEIN APBE PRECURSOR pir  C71281 conserved hypothetical protein TP0796 - syphilis spirochete gb AAC65759.1  (AE001250) conserved hypothetical protein [Treponema pallidum] Length = 362
89	44	26	54	50
Contig109 (27573-28847 m)	Contig109 (26686-27474 p)	Contig109 (25352-26587 p)	Contig109 (24726-25328 p)	Contig109 (23755-24708 p)
SA-1062.1	SA-1063.1	SA-1064.1	SA-1065.1	SA-1066.1
SeqID 201	SeqID 202	SeqID 203	SeqID 204	SeqiD 205



SeqID 206	SA-1067.1	Contig109 (23439-23621 p)	55	Identities = 27/60 (45%), Positives = 36/60 (60%) splQ01468 XYLH_PSEPU 4-OXALOCROTONATE TAUTOMERASE (4-OT) pir  A43397 4-oxalocrotonate tautomerase (EC 5.3.2) xylH - Pseudomonas putida plasmid TOL pWW0 gb AAA25694.1  (M94186) 4-oxalocrotonate tautomerase [Pseudomonas putida] gb AAA26046.1  (M95650) 4-oxalocrotonate tautomerase [Plasmid pWW0] prf  1916401D 4-oxalocrotonate tautomerase [Pseudomonas putida] Length = 63
SeqID 207	SA-1068.1	Contig109 (22732-23301 m)	89	Identities = 157/189 (83%), Positives = 175/189 (92%) spjP47848 KITH_STRGC THYMIDINE KINASE gbjAAB02289.1  (L40415) thymidine kinase [Streptococcus gordonii] Length = 191
SeqID 208	SA-1069.1	Contig109 (21618-22697 m)	76	Identities = 211/351 (60%), Positives = 281/351 (79%), Gaps = 1/351 (0%) splP45872 RF1_BACSU PEPTIDE CHAIN RELEASE FACTOR 1 (RF-1) pir  S55437 translation releasing factor RF-1 Bacillus subtilis emb CAA89884.1  (Z49782) peptide chain release factor 1 [Bacillus subtilis] emb CAB15718.1  (Z99122) peptide chain release factor 1 [Bacillus subtilis] Length = 356
SeqID 209	SA-107.1	Contig131 (15332-16198 p)	78	Identities = 195/277 (70%), Positives = 236/277 (84%) gb AAA56773.1  (U17382) putative multiple membrane domain protein; possible TTG initiation codon at position 1064, near putative RBS at position 1052 [Streptococcus pyogenes] Length = 277
SeqID 210	SA-1071.1	Contig109 (20788-21618 m)	47	Identities = 94/236 (39%), Positives = 132/236 (55%), Gaps = 12/236 (5%) dbj BAB07493.1  (AP001519) protoporphyrinogen oxidase [Bacillus halodurans] Length = 289
SeqID 211	SA-1072.1	Contig109 (20199-20795 m)	22	Identities = 71/199 (35%), Positives = 115/199 (57%), Gaps = 4/199 (2%) pir  E72324 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD35934.1 AE001752_1 (AE001752) conserved hypothetical protein [Thermotoga maritima] Length = 335



	No Hits found	Contig109 (13587-13841 m)	SA-1078.2	SeqID 217
Identities = 219/552 (39%), Positives = 337/552 (60%), Gaps = 3/552 (0%) splQ9WYC4 Y288_THEMA HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN TM0288 pir]F72396 ABC transporter, ATP-binding protein - Thermotoga maritima (strain MSB8) gb AAD35376.1 AE001710_11 (AE001710) ABC transporter, ATP-binding protein [Thermotoga maritima] Length = 598	89 9		SA-1077.1	SeqID 216
Identities = 196/570 (34%), Positives = 327/570 (56%), Gaps = 5/570 (0%) pir  E72396 ABC transporter, ATP-binding protein - Thermotoga maritima (strain MSB8) gb AAD35375.1 AE001710_10 (AE001710) ABC transporter, ATP-binding protein [Thermotoga maritima] Length = 577		Contig109 (15532-17253 m)	SA-1076.1	SeqID 215
Identities = 53/140 (37%), Positives = 78/140 (54%), Gaps = 2/140 (1%) pir  F69900 hypothetical protein yocA [imported] - Bacillus subtilis gb AAB84433.1  (AF027868) RAS-related protein [Bacillus subtilis] emb CAB13805.1  (299114) similar to transposon-related protein [Bacillus subtilis]	98	Contig109 (17265-17867 m)	SA-1075.1	SeqID 214
Identities = 32/95 (33%), Positives = 53/95 (55%), Gaps = 1/95 (1%) emb CAB71302.1  (AJ130879) hypothetical protein   [Clostridium sticklandii]	15	Contig109 (17869-18846 m)	SA-1074.1	SeqID 213
Identities = 242/417 (58%), Positives = 308/417 (73%), Gaps = 7/417 (1%) splQ9WZH9JGLYA_THEMA SERINE HYDROXYMETHYLTRANSFERASE (SERINE METHYLASE) (SHMT) pirl F72341 glycine hydroxymethyltransferase (EC 2.1.2.1) - Thermotoga maritima (strain MSB8) gb AAD35802.1 AE001743_4 (AE001743) serine hydroxymethyltransferase [Thermotoga maritima] Length = 427	73	Contig109 (18851-20107 m)	SA-1073.1	SeqID 212



SeqID 218	SA-1079.1	Contig109 (11805-13523 p)	91	Identities = 486/573 (84%), Positives = 527/573 (91%), Gaps = 1/573 (0%) embjCAB95418 1  (AJ243290) phosphoglucomutase   Streptococcus thermophilus   Length = 572
SeqID 219	SA-108.1	Contig131 (14614-15276 p)	45	Identities = 51/215 (23%). Positives = 102/215 (46%), Gaps = 9/215 (4%) gb AAD04237 1  (AF007761) MutR [Streptococcus mutans] Length = 287
SeqID 220	SA-1080.1	Contig109 (11246-11695 p)	84	Identities = 49/190 (25%), Positives = 84/190 (43%), Gaps = 33/190 (17%) emb CAA80247.1  (Z22520) membrane protein [Bacillus acidopullulyticus]
SeqID 221	SA-1081.1	Contig109 (10535-11077 p)	99	Identities = 101/145 (69%), Positives = 122/145 (83%) splQ54433IDFP_STRMU DNA/PANTOTHENATE METABOLISM FLAVOPROTEIN HOMOLOG gblAAC44502.1  (U48885) DNA/pantothenate metabolism flavoprotein [Streptococcus mutans] Length = 145
SeqID 222	SA-1082.1	Contig109 (9841-10542 p)	94	Identities = 74/232 (31%), Positives = 122/232 (51%), Gaps = 36/232 (15%) spiQ58323 DFP_METJA DNA/PANTOTHENATE METABOLISM FLAVOPROTEIN HOMOLOG pir JA64414 pantothenate metabolism flavoprotein MJ0913 - Methanococcus jannaschii gb AAB98918.1  (U67535) pantothenate metabolism flavoprotein (dfp) [Methanococcus jannaschii]
SeqID 223	SA-1083.1	Contig109 (8866-9702 m)	43	Identitles = 94/221 (42%), Positives = 134/221 (60%), Gaps = 2/221 (0%) dbj BAB07541.1  (AP001520) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 278
SeqID 224	SA-1084.1	Contig109 (7674-8873 m)	54	Identities = 162/311 (52%), Positives = 218/311 (70%), Gaps = 1/311 (0%) pir[ S49950 probable trimethylamine dehydrogenase (EC 1.5.99.7) - Mycoplasma capricolum (SGC3) (fragment) pir[ S77803 hypothetical protein MC012 - Mycoplasma capricolum emb[CAA83700.1] (Z33015) similar to trimethylamine DH [Mycoplasma capricolum]

SeqID 225	SA-1085.1	Contig109 (6691-7677 m)	. 25	Identities = 121/333 (36%), Positives = 190/333 (56%), Gaps = 12/333 (3%) dbj BAB04594.1  (AP001510) unknown conserved protein [Bacillus halodurans] Length = 335
SeqID 226	SA-1086.1	Contig109 (6330-6662 m)	14	Identities = 30/91 (32%), Positives = 53/91 (57%), Gaps = 1/91 (1%) gblAAF81345.1 AC007767_25 (AC007767) Identical to a glycine cleavage system H-protein precursor from Arabidopsis thaliana gblP25855. It contains a glycine cleavage H-protein domain PF 01597. ESTs gblR90208, gblA1994794, gblAA605324, gblN38240, gblAV533336, gblAV534187, gblAA5> Length = 166
SeqID 227	SA-1087.2	Contig109 (5519-6337 m)	35	Identities = 71/171 (41%), Positives = 97/171 (56%), Gaps = 13/171 (7%) ref[NP_054786.1  LRP16 protein [Homo saplens] gb AAF15294.1 AF202922_1 (AF202922) LRP16 [Homo saplens] gb AAH00270.1 AAH00270 (BC000270) LRP16 protein [Homo sapiens] gb AAH03188.1 AAH03188 (BC003188) LRP16 protein [Homo sapiens]
SeqID 228	SA-1088.2	Contig138 (9701-11002 m)	S S	Identities = 160/415 (38%), Positives = 251/415 (59%), Gaps = 13/415 (3%) splQ45400 PTCC_BACST PTS SYSTEM, CELLOBIOSE-SPECIFIC IIC COMPONENT (EIIC-CEL) (CELLOBIOSE-PERMEASE IIC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, C COMPONENT) pir  C49898 cellobiose phosphotransferase system cells - Bacillus stearothermophilus gb AAA17390.1  (U07818) cellobiose phosphotransferase enzyme II [Bacillus stearothermophilus] Length = 451
SeqID 229	SA-1089.1	Contig138 (11004-11324 m)	55	Identities = 46/100 (46%), Positives = 62/100 (62%) pir  D82219 PTS system, cellobiose-specific IIB component VC1281 [imported] - Vibrio cholerae (group O1 strain N16961) gb[AAF9440.1] (AE004207) PTS system, cellobiose-specific IIB component [Vibrio cholerae] Length = 101



HOSPHATE PHOSPHATE E) (IMPDH) 1.1.1.205) - 3056) inosine coccus	32 (68%) scherichia coli]	56%), Gaps = otional regulator 316	//243 (60%), Gaps = //YPOTHETICAL IN OSMB-RNB Shypothetical protein (AE000226) putative (Escherichia coli K12)	/289 (49%), Gaps = FATIVE PYRUVATE YME pir  H64819 nzyme (EC 1.97.1.4) (AE000184) putative [Escherichia dependent pyruvate [Escherichia coli]
Identities = 461/493 (93%), Positives = 478/493 (96%) splP50099 IMDH_STRPY INOSINE-5 -MONOPHOSPHATE DEHYDROGENASE (IMP DEHYDROGENASE) (IMPDH) (IMPD) pir JJC4372 IMP dehydrogenase (EC 1.1.205) - Streptococcus pyogenes gb AAB03846.1  (U26056) inosine monophosphate dehydrogenase [Streptococcus	Identities = 42/102 (41%), Positives = 70/102 (68%) gb AAA23551.1  (M93570) PTS enzyme III cel [Escherichia coli] Length = 116	Identities = 98/309 (31%), Positives = 178/309 (56%), Gaps = 1/309 (0%) dbj BAB04499.1  (AP001509) transcriptional regulator [Bacillus halodurans]	Identities = 96/243 (39%), Positives = 148/243 (60%), Gaps = 1/243 (0%) splP76034 YCIT_ECOL! HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN OSMB-RNB INTERGENIC REGION pir  G64876 hypothetical protein b1284 - Escherichia coli gb AAC74366.1  (AE000226) putative DEOR-type transcriptional regulator [Escherichia coli K12 Length = 249	Identities = 106/289 (36%), Positives = 146/289 (49%), Gaps = 50/289 (17%) splP75794 PFLE_ECOLI PUTATIVE PYRUVATE FORMATE-LYASE 3 ACTIVATING ENZYME pir  H64819 probable pyruvate formate-lyase activating enzyme (EC 1.97.1.4) 3 - Escherichia coli gb AAC73911.1  (AE000184) putative pyruvate formate-lyase 2 activating enzyme [Escherichia coli K12] db BAA35512.1  (D90720) Iron-dependent pyruvate formate-lyase-activating enzyme [Escherichia coli] Length = 308
66	09	53	56	54
Contig131 (13055-14539 p)	Contig138 (11341-11670 m)	Contig138 (11868-12857 m)	Contig138 (12970-13725 m)	Contig138 (13846-14622 p)
SA-109.1	SA-1090.1	SA-1091.2	SA-1092.2	SA-1093.1
SeqID 230	SeqID 231	SeqID 232	SeqID 233	SeqID 234



SeqID 235	SA-1094.1	Contig138 (14669-15025 m)	26	Identities = 45/120 (38%), Positives = 71/120 (58%), Gaps = 11/120 (9%) dbjiBAB07066 1j (AP001518) polyribonucleotide nucleotidyltransferase (general stress protein 13) [Bacillus halodurans] Length = 138
SeqID 236	SA-1095.1	Contig138 (15027-16385 m)	. 23	Identities = 81/174 (46%). Positives = 109/174 (62%), Gaps = 30/174 (17%) splP87051 YDJ3_SCHPO PROBABLE PEPTIDYL-PROLYL CIS-TRANS ISOMERASE C57A10.03 pir  T38930 peptidylprolyl isomerase (EC 5.2.1.8) SPAC57A10.03 [similarity] - fission yeast (Schizosaccharomyces pombe) emb CAB08166.1  (294864) putative peptidyl-prolyl cis-trans isomerase [Schizosaccharomyces pombe]
SeqID 237	SA-1096.1	Contig138 (16469-17122 m)	. 83	Identities = 155/209 (74%), Positives = 184/209 (87%)   emb CAB54571.1  (AJ006393) response regulator [Streptococcus   pneumoniae]
SeqID 238	SA-1097.1	Contig138 (17103-18122 m)	70	Identities = 159/334 (47%), Positives = 239/334 (70%), Gaps = 5/334 (1%) emb CAB54570.1  (AJ006393) histidine kinase [Streptococcus pneumoniae]
SeqID 239	SA-1098.1	Contig138 (18119-18814 m)	43	Identities = 53/230 (23%), Positives = 104/230 (45%), Gaps = 14/230 (6%) dbj BAB03323.1  (AB035448) hypothetical protein [Staphylococcus aureus]
SeqID 240	SA-1099.1	Contig138 (18989-19231 p)	No Hits found	
SeqID 241	SA-11.1	Contig137 (32951-36100 p)	27	Identities = 170/605 (28%), Positives = 289/605 (47%), Gaps = 105/605 (17%) ref[NP_05317.1] pXO2-16 [Bacillus anthracis] gb AAF13621.1 AF188935_19 (AF188935) pXO2-16 [Bacillus anthracis]
SeqID 242	SA-110.1	Contig131 (12416-12901 p)	56	Identities = 63/144 (43%), Positives = 93/144 (63%), Gaps = 4/144 (2%) emb CAA09426.1  (AJ010954) arginine repressor [Bacillus stearothermophilus] Length = 149

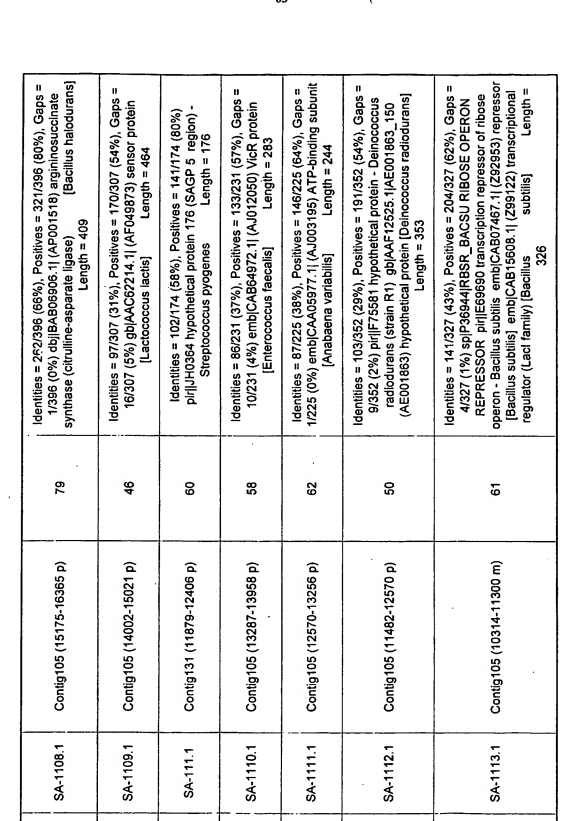


SealO 243	SA-1100 1	Contict 38 (18973-20928 m)	43	Identities = 219/380 (57%), Positives = 284/380 (74%), Gaps = 8/380 (2%) emblCA410713 11 (A.1132604) hypothetical protein
410 643		T.	?	[Lactococcus lactis] Length = 378
SeqID 244	SA-1101.1	Contig138 (20928-21665 m)	69	Identities = 131/245 (53%), Positives = 177/245 (71%), Gaps = 4/245 (1%) emb CAA10712.1  (AJ132604) pppL protein [Lactococcus lactis] Length = 258
SeqID 245	SA-1102.1	Contig138 (21703-23025 m)	89	Identities = 222/434 (51%), Positives = 305/434 (70%), Gaps = 15/434 (3%) emb[CAA10711.1  (AJ132604) sunL protein [Lactococcus lactis] Length = 424
SeqID 246	SA-1103.1	Contig138 (23015-23950 m)	. 29	Identities = 158/314 (50%), Positives = 215/314 (68%), Gaps = 5/314 (1%) dbj BAB06227.1  (AP001515) methionyl-tRNA formyltransferase [Bacillus halodurans] Length = 317
SeqID 247	SA-1104.1	Contig138 (23997-26387 m)	99	Identities = 377/807 (46%), Positives = 529/807 (64%), Gaps = 21/807 (2%) splP94461 PRIA_BACSU PRIMOSOMAL PROTEIN N (REPLICATION FACTOR Y) pir  A69682 primosomal replication factor Y priA - Bacillus subtilis emb CAA74261.1  (Y13937) putative PriA protein [Bacillus subtilis] emb CAB13444.1  (Z99112) primosomal replication factor Y (primosomal protein N%) [Bacillus subtilis] Length = 805
SeqID 248	SA-1105.2	Contig138 (26461-26775 m)	35	Identities = 25/57 (43%), Positives = 40/57 (69%), Gaps = 1/57 (1%) pir  C69878 hypothetical protein yloH - Bacillus subtilis emb CAA74272.1  (Y13937) putative rpoZ protein [Bacillus subtilis] emb CAB13442.1  (Z99112) yloH [Bacillus subtilis] Length = 67
SeqID 249	SA-1107.1	Contig105 (16384-17661 p)	75	Identities = 260/416 (62%), Positives = 324/416 (77%)   dbj BAB06905.1  (AP001518) argininosuccinate lyase [Bacilius halodurans]   Length = 458

SeqID 256

SeqID 255





SeqID 250

SeqID 252

SeqID 251

SeqID 253

SeqID 254



Identities = 132/293 (45%), Positives = 176/293 (60%), Gaps = 4/293 (1%) splP36945 RBSK_BACSU RIBOKINASE pir  D69690 ribokinase (EC 2.7.1.15) - Bacillus subtilis emb CAB07465.1  (292953) ribokinase [Bacillus subtilis] emb CAB15609.1  (299122) ribokinase [Bacillus subtilis] Length = 293	Identities = 74/131 (56%), Positives = 96/131 (72%), Gaps = 1/131 (0%) splP36946 RBSD_BACSU HIGH AFFINITY RIBOSE TRANSPORT PROTEIN RBSD pir  140464 ribose ABC transporter (membrane protein) rbsD - Bacillus subtilis emb CAA81050.1  (225798) Membrane Ribose Binding Protein [Bacillus subtilis] emb CAB15610.1  (299122) ribose ABC transporter (membrane protein) [Bacillus subtilis]	Identities = 298/493 (60%), Positives = 377/493 (76%), Gaps = 1/493 (0%) splP36947 RBSA_BACSU RIBOSE TRANSPORT ATP-BINDING PROTEIN RBSA pir  H69689 ribose ABC transporter (ATP-binding protein) rbsA - Bacillus subtilis emb CAB07463.1  (Z92953) ATP-binding transport protein [Bacillus subtilis] emb CAB15611.1  (Z99122) ribose ABC transporter (ATP-binding protein) [Bacillus subtilis]	Identities = 198/305 (64%), Positives = 252/305 (81%), Gaps = 1/305 (0%) pir  B69690 ribose ABC transporter (permease) rbsC - Bacillus subtilis emb CAB07462.1  (Z92953) membrane transport protein [Bacillus subtilis] emb CAB15612.1  (Z99122) ribose ABC transporter (permease) [Bacillus subtilis]
57	09	. 9/	. 82
Contig105 (9410-10321 m)	Contig105 (9037-9435 m)	Contig105 (7543-9021 m)	Contig105 (6600-7541 m)
SA-1114.1	SA-1115.1	SA-11:16.1	SA-1118.1
SeqID 257	SeqID 258	SeqID 259	SeqID 260

SA-1121.1

SeqID 264



		65 .	
Identities = 143/301 (47%), Positives = 204/301 (67%), Gaps = 1/301 (0%) splP36949 RBSB_BACSU D-RIBOSE-BINDING PROTEIN PRECURSOR pir A69690 ribose ABC transporter (ribose-binding protein) rbsB - Bacillus subtilis emb CAB07461.1  (Z92953) periplasmic substrate-binding protein [Bacillus subtilis] emb CAB15613.1  (Z99122) ribose ABC transporter (ribose-binding protein) [Bacillus subtilis]	Identities = 42/132 (31%), Positives = 74/132 (55%), Gaps = 3/132 (2%) pir  H83123 hypothetical protein PA4181 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG07568.1 AE004834_8 (AE004834) hypothetical protein [Pseudomonas aeruginosa] Length = 239	Identities = 273/491 (55%), Positives = 353/491 (71%), Gaps = 19/491 (3%) splP22250 SYE_BACSU GLUTAMYL-TRNA SYNTHETASE (GLUTAMATETRNA LIGASE) (GLURS) pir  SYBSET glutamatetRNA ligase (EC 6.1.17) gltX - Bacillus subtilis gb AAA22495.1  (M55073) transfer RNA-Gln synthetase [Bacillus subtilis] gb AAA21796.1  (14580) glutamyl-tRNA synthetase [Bacillus subtilis] db  BAA05326.1  (D26185) glutamyl-tRNA synthetase [Bacillus subtilis] emb CAB11868.1  (299104) glutamyl-tRNA synthetase [Bacillus subtilis] Length = 483	Identities = 201/443 (45%), Positives = 290/443 (65%) splP77212 YKGC_ECOLI PROBABLE PYRIDINE NUCLEOTIDE- DISULFIDE OXIDOREDUCTASE IN EAEH-BETA INTERGENIC REGION pir H64756 probable mercury(II) reductase (EC 1.16.1.1) - Escherichia coli gb AAB18031.1  (U73857) similar to S. aureus mercury(II) reductase [Escherichia coli] gb AAC73407.1  (AE000137) putative oxidoreductase [Escherichia coli K12] Length = 450
. 69	31	69	,
Contig 105 (5579-6547 m)	Contig131 (10814-11482 m)	Contig105 (4085-5539 p)	Contig105 (2557-3876 p)

SA-1120.1

SeqID 263

SA-1119.1

SeqID 261

SeqID 262



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Identities = 69/162 (42%), Positives = 100/162 (61%), Gaps = 2/162 (1%) splQ10612 YC84_MYCTU HYPOTHETICAL 18.2 KDA PROTEIN RV1284 pir  H70771 hypothetical protein Rv1284 - Mycobacterium tuberculosis (strain H37RV) emb CAA97750.1  (Z73419) hypothetical protein Rv1284 [Mycobacterium tuberculosis] Length = 163	Identities = 284/453 (62%), Positives = 358/453 (78%), Gaps = 4/453 (0%) sp[P37572 RADA_BACSU DNA REPAIR PROTEIN RADA HOMOLOG (DNA REPAIR PROTEIN SMS HOMOLOG) pir[IS66116 DNA repair protein sms - Bacillus subtilis dbj[BAA05321.1] (D26185) unknown [Bacillus subtilis] emb[CAB11863.1] (299104) DNA repair protein homolog [Bacillus subtilis]	Identities = 41/80 (51%), Positives = 52/80 (64%), Gaps = 5/80 (6%) gb AAF74088.1  (AF212845) putative dUTPase [Lactococcus lactis bacterlophage ul36]	Identities = 46/128 (35%), Positives = 70/128 (53%), Gaps = 5/128 (3%) gb AAK05804. i AE006401_5 (AE006401) mannose-specific PTS system component IIAB (EC 2.7.1.69) [Lactococcus lactis subsp. lactis] Length = 329	Identities = 156/370 (42%), Positives = 219/370 (59%), Gaps = 3/370 (0%) dbj BAB05773.1  (AP001514) unsaturated glucuronyi hydrolase [Bacillus halodurans] Length = 375	Identities = 63/125 (50%), Positives = 89/125 (70%), Gaps = 1/125 (0%) gblAAC44679.1  (U65015) PTS permease for mannose subunit IIIMan C terminal domain [Vibrio furnissii] Length = 157	Identities = 93/295 (31%), Positives = 140/295 (46%), Gaps = 48/295 (16%) gb AAF81084.1 AF228498_4 (AF228498) AgaW [Escherichia coli] Length = 259
99	78	64	44	54	52	45
Contig105 (1906-2403 p)	Contig105 (406-1770 p)	Contig105 (2-244 p)	Contig120 (17546-17980 p)	Contig120 (17983-19179 p)	Contig120 (19234-19728 p)	Contig120 (19764-20630 p)
SA-1122.1	SA-1123.1	SA-1124.1	SA-1125.2	SA-1126.2	SA-1127.1	SA-1129.1
SeqID 265	SeqID 266	SeqID 267	SeqID 268	SeqID 269	SeqID 270	SeqID 271





SeqID 272	SA-113.1	Contig131 (9315-10547 m)	98	Identities = 375/411 (91%), Positives = 395/411 (95%), Gaps = 1/411 (0%) sp P16962 ARCA_STRPY ARGININE DEIMINASE (ADI) (ARGININE DIHYDROLASE) (AD) (STREPTOCOCCAL ACID GLYCOPROTEIN) pir  A38835 streptococcal antitumor protein - Streptococcus pyogenes emb CAA39192.1  (X55659) antitumor protein (Streptococcus pyogenes) db  BAA02938.1  (D13790) acid glycoprotein (Streptococcus pyogenes]
SeqID 273	SA-1131.1	Contig120 (20617-21432 p)	54	Identities = 101/278 (36%), Positives = 164/278 (58%), Gaps = 6/278 (2%) gb AAA57943.1  (U18997) ORF_0290; Geneplot suggests frameshift linking to 0267, not found [Escherichia coli] Length = 290
SeqID 274	SA-1132.1	Contig120 (21512-23416 p)	6	Identities = 76/206 (36%), Positives = 124/206 (59%), Gaps = 1/206 (0%) emb CAB01924.1  (Z79691) OrfA [Streptococcus pneumoniae] Length = 207
SeqID 275	SA-1134.1	Contig120 (23496-24497 p)	79	Identities = 222/333 (66%), Positives = 279/333 (83%) emb[CAB01925.1  (Z79691) RegR [Streptococcus pneumoniae] Length = 333
SeqID 276	SA-1135.1	Contig120 (24568-25182 m)	. 28	Identities = 169/204 (82%), Positives = 192/204 (93%), Gaps = 1/204 (0%) emb CAC15392.1  (AJ278785) polypeptide deformylase [Streptococcus pneumoniae] Length = 203
SeqID 277	SA-1136.1	Contig120 (25248-25760 m)	42	Identities = 58/191 (30%), Positives = 98/191 (50%) sp[P33023]YEIL_ECOLI HYPOTHETICAL 25.3 KD PROTEIN IN NFO-FRUA INTERGENIC REGION pir]B64985 hypothetical 25.3 kD protein in nfo-fruA intergenic region - Escherichia coli (strain K-12) gb[AA460515.1] (U00007) yeil [Escherichia coli] gb[AAC75224.1] (AE000305) putative transcriptional regulator [Escherichia coli K12] prf[ 20142538J yeil gene [Escherichia coli] Length = 219



, Gaps = 24B0761 - 30057.1	), Gaps = ported] - ) putative Length =	38%)	3%) PROTEIN protein nes	, Gaps = SAL 36.4 SION - Bacillus subtilis]		(52%), Gaps = Il protein yvgP - I) similar to Length = 670
Identities = 95/382 (24%), Positives = 174/382 (44%), Gaps = 30/382 (7%) pir[ID75094 transport protein, permease PAB0761 Pyrococcus abyssi (strain Orsay) emb[CAB50057.1  (AJ248286) TRANSPORT PROTEIN, permease [Pyrococcus abyssi] Length = 372	Identities = 101/327 (30%), Positives = 169/327 (50%), Gaps = 12/327 (3%) pir  T50594 probable oxidoreductase [imported] - Streptomyces coelicolor emb CAB61731.1  (AL133220) putative oxidoreductase. [Streptomyces coelicolor A3(2)] Length = 351	Identities = 504/631 (79%), Positives = 563/631 (88%) emb CAC14579.1  (AJ249396) oligopeptidase [Streptococcus thermophilus]	Identities = 66/114 (57%), Positives = 85/114 (73%) splP16963 YSA1_STRPY HYPOTHETICAL 16.6 KD PROTEIN IN SAGP 3 REGION pir  JE0061 hypothetical 16.5K protein (SAGP 5 region) - Streptococcus pyogenes dbj BAA02939.1  (D13790) ORF2 (Streptococcus pyogenes) Length = 146	Identities = 133/309 (43%), Positives = 197/309 (63%), Gaps = 5/309 (1%) splO35046 YOCD_BACSU HYPOTHETICAL 36.4 KDA PROTEIN IN CSA4-DES INTERGENIC REGION pir  A69901 immunity to bacteriotoxins homolog yocD - Bacillus subtilis gblAAB84435.1  (AF027868) YocD [Bacillus subtilis] emb CAB13809.1  (Z99114) similar to immunity to bacteriotoxins [Bacillus subtilis]		Identities = 174/524 (33%), Positives = 275/524 (52%), Gaps = 13/524 (2%) pir  E70040 conserved hypothetical protein yvgP - Bacillus subtilis emb CAB15347.1  (299121) similar to hypothetical proteins [Bacillus subtilis]
43	20	88	25		No Hits found	68
Contig120 (26052-27167 p)	Contig120 (27213-28196 m)	Contig120 (28344-30239 p)	Contig131 (8878-9219 m)	Contig120 (30289-31242 m)	Contig120 (31400-31645 p)	Contig120 (31746-33716 p)
SA-1137.1	SA-1138.1	SA-1139.1	SA-114.1	SA-1140.1	SA-1141.1	SA-1143.2
SeqID 278	SeqID 279	SeqID 280	SeqID 281	SeqID 282	SeqID 283	SeqID 284

Identities = 383/509 (75%), Positives = 441/509 (86%) dbj BAA25696.1  (AB010712) NADH oxidase/alkyl hydroperoxidase reductase Length = 510	dentities = 168/186 (90%), Positives = 180/186 (96%)  dbj BAA25695.1  (AB010712) alkyl hydroperoxidase  Streptococcus mutans]	Identities = 36/72 (50%), Positives = 49/72 (68%) splP54457 YQEL_BACSU HYPOTHETICAL 13.3 KD PROTEIN IN AROD-COMER INTERGENIC REGION pir] H69951 ybeB protein homolog yqeL - Bacillus subtilis dbj BAA12449.1  (D84432) YqeL [Bacillus subtilis] emb CAB14504.1  (Z99117) similar to hypothetical proteins [Bacillus subtilis]	Identities = 86/242 (35%), Positives = 155/242 (63%), Gaps = 4/242 (1%) splP54458 YQEM_BACSU HYPOTHETICAL 28.3 KD PROTEIN IN AROD-COMER INTERGENIC REGION pir  A69952 conserved hypothetical protein yqeM - Bacillus subtilis dbj BAA12450.1  (D84432) YqeM [Bacillus subtilis] emb CAB14503.1  (299117) similar to hypothetical proteins [Bacillus subtilis]		Identities = 244/333 (73%), Positives = 294/333 (88%), Gaps = 1/333 (0%) pir  T46742 ornithine carbamoyltransferase (EC 2.1.3.3) [validated] - Lactobacillus sakei emb CAA04683.1  (AJ001330) ornithine transcarbamoylase [Lactobacillus sakei] Length = 337	Identities = 131/367 (35%), Positives = 185/367 (49%), Gaps = 45/367 (12%) dbj BAB06304.1  (AP001516) unknown conserved protein [Bacillus halodurans] Length = 416
86	26	67	6	No Hits found	86	49
Contig120 (33993-35525 m)	Contig120 (35543-36103 m)	Contig124 (43925-44143 m)	Contig124 (43125-43859 m)	Contig124 (42851-43075 m)	Contig131 (7849-8862 m)	Contig124 (41745-42854 m)
SA-1144.1	SA-1145.1	SA-1146.1	SA-1148.1	SA-1149.1	SA-115.1	SA-1150.1
SeqID 285	SeqID 286	SeqID 287	SeqiD 288	SeqID 289	SeqID 290	SeqID 291



Identities = 265/470 (56%), Positives = 347/470 (73%), Gaps = 3/470 (0%) emb CAA76779.1  (Y17554) permease [Bacillus licheniformis] Length = 468	Identities = 182/237 (76%), Positives = 201/237 (84%) splP76351 YEEN_ECOL! HYPOTHETICAL 25.9 KDA PROTEIN IN AMN-CBL INTERGENIC REGION pir  A64963 conserved hypothetical protein b1983 - Escherichia coli gb AAC75047.1  (AE000290) orf, hypothetical protein [Escherichia coli K12] Length = 238	Identities = 72/178 (40%), Positives = 101/178 (56%), Gaps = 2/178 (1%) pir  T36850 hypothetical protein SCI35.37 - Streptomyces coelicolor emb CAA20826.1  (AL031541) hypothetical protein SCI35.37 [Streptomyces coelicolor A3(2)] Length = 242	Identities = 83/264 (31%), Positives = 133/264 (49%), Gaps = 13/264 (4%) gb[AAC45332.1  (U97348) basic surface protein [Lactobacillus fermentum] Length = 264	Identities = 110/284 (38%), Positives = 176/284 (61%), Gaps = 13/284 (4%) emb CAB59825.1  (AJ012388) hypothetical protein [Lactococcus lactis]	Identities = 132/419 (31%), Positives = 212/419 (50%), Gaps = 14/419 (3%) pir  E75327 ArgE/DapE/Acy1 family protein - Deinococcus radiodurans (strain R1) gb AAF11560.1 AE002038_9 (AE002038) ArgE/DapE/Acy1 family protein [Deinococcus radiodurans] Length = 463	Identities = 43/115 (37%), Positives = 52/115 (44%), Gaps = 2/115 (1%) pir[]H72617 hypothetical protein APE1401 - Aeropyrum pernix (strain K1) dbj[BAA80398.1  (AP000061) 123aa long hypothetical protein [Aeropyrum pernix] Length = 123
72	72	43	46	58	46	88
Contig131 (6359-7786 m)	Contig124 (34888-35604 m)	Contig124 (34044-34733 m)	Contig124 (33109-33942 m)	Contig124 (32144-32968 m)	Contig124 (30634-32010 m)	Contig124 (29833-30207 p)
SA-116.1	SA-1160.1	SA-1161.1	SA-1162.1	SA-1163.1	SA-1165.1	SA-1166.1
SeqID 300	SeqID 301	SeqID 302	SeqID 303	SeqID 304	SeqID 305	SeqID 306



SeqID 307	SA-1167.1	Contig124 (29571-30641 m)	12	Identities = 187/338 (55%), Positives = 256/338 (75%), Gaps = 12/338 (3%) emb[CAB59828 1] (AJ012388) hypothetical protein [Lactococcus lactis]
SeqID 308	SA-1169.1	Contig124 (28882-29574 m)	73	Identities = 137/231 (59%), Positives = 171/231 (73%), Gaps = 1/231 (0%) emb[CAB59829 1] (AJ012388) hypothetical protein [Lactococcus lactis] Length = 231
SeqID 309	SA-117.1	Contig131 (5382-6338 m)	76	Identities = 199/311 (63%), Positives = 254/311 (80%), Gaps = 3/311 (0%) splO53090JARCC_LACSK CARBAMATE KINASE pir  T46743 carbamate kinase (EC 2.7.2.2) [validated] - Lactobacillus sakei emb CAA04684.1  (AJ001330) carbamate kinase [Lactobacillus sakei]
SeqID 310	SA-1170.1	Contig124 (27778-28833 p)	72	Identities = 196/347 (56%), Positives = 257/347 (73%), Gaps = 1/347 (0%) pir  C81088 alcohol dehydrogenase, zinc-containing NMB1395 [imported] - Neisseria meningitidis (group B strain MD58) gb AAF41759.1  (AE002488) alcohol dehydrogenase, zinc-containing (Neisseria meningitidis MC58] Length = 346
SeqID 311	SA-1171.2	Contig124 (26294-27664 p)	55	Identities = 182/453 (40%), Positives = 265/453 (58%), Gaps = 29/453 (6%) pir] C69596 branched-chain amino acid transporter braB - Bacillus subtilis gb AAC00400.1  (AF008220) branch-chain amino acid transporter [Bacillus subtilis] emb CAB14938.1  (Z99119) branched-chain amino acid transporter [Bacillus subtilis]
SeqID 312	SA-1173.3	Contig136 (3489-3797 m)	No Hits found	
SeqID 313	SA-1174.1	1~	No Hits found	
SeqID 314	SA-1175.1	Contig136 (4104-4676 m)	No Hits found	
SeqID 315	SA-1176.1	_	No Hits found	
SeqID 316	SA-1177.1	)	No Hits found	
SeqID 317	SA-1178.1	$\sim$	No Hits found	
SeqID 318	SA-1179.1	Contig136 (6986-7477 m)	No Hits found	

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Identities = 520/568 (91%), Positives = 547/568 (95%), Gaps = 1/568 (0%) splP23531 PTLB_LACLA PTS SYSTEM, LACTOSE-SPECIFIC IIBC COMPONENT (EIIBC-LAC) (LACTOSE-ENZYME II, BC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, BC COMPONENT) (EII-LAC) pirl B23696 phosphotransferase system enzyme II (EC 2.7.1.69) - Lactococcus lactis gb AAA25182.1  (M60447) enzyme III [Lactococcus lactis]		Identities = 89/105 (84%), Positives = 97/105 (91%) splP23532 PTLA_LACLA PTS SYSTEM, LACTOSE-SPECIFIC IIA COMPONENT (EIIA-LAC) (LACTOSE-PERMEASE IIA COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, A COMPONENT) (FIII-LAC) pir  A23696 phosphotransferase system enzyme III (EC 2.7) - Lactococcus lactis pdb 1E2A A Chain A, Enzyme Iia From The Lactose Specific Pts From Lactococcus Lactis pdb 1E2A B Chain B, Enzyme Iia From The Lactose Specific Pts From Lactococcus Specific Pts From Lactococcus Specific Pts From Lactose Specific Pts From Lactococcus Specific Pts From Lactose Specific Pts From Lactose Specific Pts From Lactose Specific Pts From Lactococcus Lactis gb AAA25181.1  (M60447) enzyme III [Lactococcus lactis] Length = 105	Identities = 104/278 (37%), Positives = 169/278 (60%), Gaps = 1/278 (0%) sp P24401 LACT_LACCA TRANSCRIPTION ANTITERMINATOR LACT emb CAB02555.1 (280834) LacT [Lactobacillus casei] gb AAB49331.1 (U21391) LacT [Lactobacillus casei]
76	No Hits found	. 78	99
Contig136 (16094-17800 m)	Contig131 (4008-4139 p)	Contig136 (17800-18117 m)	Contig136 (18146-18979 m)
SA-1189.1	SA-119.1	SA-1190.1	SA-1191.1
SeqiD 326	SeqID 327	SeqiD 328	SeqID 329

pir[[039778] pir[[039778] ccus lactis lactis] dolase	1%), Gaps = OSE-6-INASE) 1) LacC - Lactococcus kinase	1 (91%) SPHATE Ictose-6- LacB - Lactococcus somerase	1 (94%) SPHATE Ictose-6- LacA - [Lactococcus Somerase	
spiP26593 LACD_LACLA TAGATOSE 1,6-DIPHOSPHATE ALDOLASE (TAGATOSE-BISPHOSPHATE ALDOLASE) (D-TAGATOSE-1,6-BISPHOSPHATE ALDOLASE) LacD tagatose-1,6-diphosphate aldolase - Lactococcus lactis gb AAA25171.1  (M65190) lacD [Lactococcus lactis] gb AAA25180.1  (M60447) tagatose 1,6-diP aldolase [Lactococcus lactis] Lactococcus lactis]	Identities = 207/310 (66%), Positives = 245/310 (78%), Gaps = 1/310 (0%) sp P23391 LACC_LACLA TAGATOSE-6-PHOSPHATE KINASE (PHOSPHOTAGATOKINASE) pir  C39778 tagatose-6-phosphate kinase (EC 2.7.1) LacC-Lactococcus lactis gb AAA25170.1  (M65190) lacC [Lactococcus lactis] gb AAA25179.1  (M60447) tagatose 6-P kinase [Lactococcus lactis]	Identities = 142/171 (83%), Positives = 156/171 (91%) splP23495 LACB_LACLA GALACTOSE-6-PHOSPHATE ISOMERASE LACB SUBUNIT pir  B39778 galactose-6-phosphate isomerase (EC 5.3.1) component LacB - Lactococcus lactis gb AAA25169.1  (M65190) lacB [Lactococcus lactis] gb AAA25178.1  (M60447) galactose 6-P isomerase [Lactococcus lactis]	Identities = 120/141 (85%), Positives = 134/141 (94%) splP23494 LACA_LACLA GALACTOSE-6-PHOSPHATE ISOMERASE LACA SUBUNIT pir  A39778 galactose-6-phosphate isomerase (EC 5.3.1) component LacA - Lactococcus lactis gb AAA25168.1  (M65190) lacA [Lactococcus lactis] gb AAA25177.1  (M60447) galactose 6-P isomerase [Lactococcus lactis]	
	78	85	76	No Hits found
Contig136 (19372-20352 m)	Contig136 (20357-21286 m)	Contig136 (21299-21814 m)	Contig136 (21831-22256 m)	Contig111 (10300-10728 p)
SA-1192.1	SA-1193.1	SA-1194.3	SA-1195.3	SA-1196.2
SeqID 330	SeqID 331	SeqID 332	SeqID 333	SeqID 334



TEIN   76)						3/89	ps = 8538) 94		os = reus]						Sillus
Identities = 35/91 (38%), Positives = 51/91 (55%)   -sp P34159 YHS1_CLOAB HYPOTHETICAL 11.0 KD PROTEIN IN HSP18 3 REGION (ORFA1) emb CAA46375.1  (X65276)   ORFA1 [Clostridium acetobutylicum]   Length = 96						Identities = 36/89 (40%), Positives = 52/89 (57%), Gaps = 3/89 (3%) pir  T39903 serine-rich protein - fission yeast (Schizosaccharomyces pombe) emb CAA22127.1  (AL033534) hypothetical serine-rich secreted protein [Schizosaccharomyces pombe] Length = 534	Identities = 171/402 (42%), Positives = 248/402 (61%), Gaps = 46/402 (11%) pir  A33952 58K mobilization protein - Streptococcus pneumoniae plasmids gb AAA25387.1  (M28538) mobilization peptide [Plasmid pMV158] Length = 494		Identities = 90/264 (34%), Positives = 139/264 (52%), Gaps = 22/264 (8%) prf[1405330A repM gene [Staphylococcus aureus] Length = 314		Identities = 34/102 (33%), Positives = 56/102 (54%) gb[AAD41248.1]AF106927_1 (AF106927) unknown [Streptococcus suis] Length = 272	•			Identities = 23/63 (36%), Positives = 36/63 (56%) dbj BAB05162.1  (AP001512) transcriptional regulator [Bacillus halodurans]
9	No Hits found	34	59	No Hits found		No Hits found	49	No Hits found	No Hits found	No Hits found	15				
Contig111 (9929-10219 p)	Contig111 (9134-9421 m)	Contig111 (8809-9144 m)	Contig110 (1487-1594 p)	Contig111 (8448-8687 m)	Contig111 (8034-8504 p)	Contig111 (6984-7439 m)	Contig111 (6610-7866 p)	Contig111 (5859-6293 p)	Contig111 (4967-5755 p)	Contig111 (4026-4667 p)		Contig131 (3142-3447 m)	Contig111 (3409-3690 p)	Contig111 (2975-3268 p)	Contig111 (2259-2810 m)
SA-1197.1	SA-1198.1	SA-1199.1	SA-120.1	SA-1200.1	SA-1201.1	SA-1202.1	SA-1203.1	SA-1205.1	SA-1207.1	SA-1208.1	SA-1209.1	SA-121.1	SA-1210.1	SA-1211.1	SA-1212.1
SeqID 335	SeqID 336	SeqID 337	SeqID 338	SeqID 339	SeqID 340	SeqID 341	SeqID 342	SeqID 343	SeqID 344	SeqID 345	SeqID 346	SeqID 347	SeqID 348	SeqID 349	SeqID 350

SeqID 351	SA-1213.1	Contig111 (1036-2190 m)	23	Identities = 128/386 (33%), Positives = 208/386 (53%), Gaps = 18/386 (4%) gb AAG29618:1  (AF217235) integrase-like protein [Staphylococcus aureus] Length = 378
SeqID 352	SA-1214.1	Contig111 (516-908 p)	£2	Identities = 90/129 (69%), Positives = 106/129 (81%) sp P07842 RS9_BACST 30S RIBOSOMAL PROTEIN S9 (BS10) pir  R3BS9 ribosomal protein S9 - Bacillus stearothermophilus Length = 129
SeqID 353	SA-1215.1	Contig111 (49-495 p)	89	Identities = 89/144 (61%), Positives = 113/144 (77%) dbj BAB03887.1  (AP001507) ribosomal protein L13 [Bacillus halodurans]
SeqID 354	SA-1216.2	Contig130 (62901-63572 p)	24	Identities = 27/119 (22%), Positives = 54/119 (44%), Gaps = 10/119 (8%) emb CAC12089.1  (AL445066) NADH dehydrogenase, chain M related protein acidophilum]
SeqID 355	SA-1217.1	Contig130 (62156-62926 p)	99	Identities = 121/249 (48%), Positives = 172/249 (68%) pir  H70023 N-acetyl-glucosamine catabolism homolog yutF - Bacillus subtilis emb CAB15219.1  (Z99120) similar to N-acetyl- glucosamine catabolism [Bacillus subtilis] Length = 256
SeqID 356	SA-1218.1	Contig130 (61418-62155 p)	42	Identities = 59/248 (23%), Positives = 104/248 (41%), Gaps = 30/248 (12%) pir  S51698 oleoyl-[acyl-carrier-protein] hydrolase (EC 3.1.2.14) - Arabidopsis thaliana pir  S69197 oleoyl-[acyl-carrier-protein] hydrolase (EC 3.1.2.14) (clone TE 1-7) - Arabidopsis thaliana emb CAA85389.1  (Z36912) acyl-(acyl-carrier protein) thioesterase [Arabidopsis thaliana] dbj BAB02069.1  (AB026647) acyl carrier protein thioesterase [Arabidopsis 1362
SeqID 357	SA-1219.1	Contig130 (60284-61414 p)	65	Identities = 173/375 (46%), Positives = 248/375 (66%), Gaps = 5/375 (1%) dbjjBAB05062.1I (AP001511) coproporphyrinogen III oxidase [Bacilius halodurans] Length = 385
SeqID 358	SA-1220.1	Contig130 (59812-60192 p)	No Hits found	



	· · ·		1
Identities = 20/68 (29%), Positives = 35/68 (51%) sp P24252 YBGA_ECOLI HYPOTHETICAL 20.2 KDA PROTEIN IN RHSC-PHRB INTERGENIC REGION (TKP) (ORF169) pir  B64806 ybgA protein - Escherichia coli gb AAA24387.1  (K01299) unidentified protein ORF169 [Escherichia coli] gb AAC73801.1  (AE000174) orf, hypothetical protein [Escherichia coli K12] dbj BAA35366.1  (D90709) Hypothetical coli dbj BAA35371.1  (D90710) Hypothetical 20.2 kd protein in phrB 5 region (tkp) (orf169). [Escherichia coli] bhrB 5 region (tkp) (orf169). [Escherichia coli]	Identities = 284/451 (62%), Positives = 353/451 (77%), Gaps = 4/451 (0%) pir  B69745 phosphoglucomutase (glycolysis) homolog ybbT - Bacillus subtilis emb CAB11953.1  (Z99104) similar to phosphoglucomutase (glycolysis) [Bacillus subtilis] emb CAB11970.1  (Z99105) similar to phosphoglucomutase (glycolysis) [Bacillus Length = 448	Identities = 90/324 (27%), Positives = 167/324 (50%), Gaps = 18/324 (5%) pir] A69745-hypothetical protein ybbR - Bacillus subtilis emb CAB11952.1  (299104) ybbR [Bacillus subtilis] emb CAB11969.1  (299105) ybbR [Bacillus subtilis] Length = 483	Identities = 125/253 (49%), Positives = 186/253 (73%), Gaps = 5/253 (1%) pir  H69744 conserved hypothetical protein ybbP - Bacillus subtilis dbj BAA19509.1  (AB002150) YbbP [Bacillus subtilis] emb CAB11951.1  (Z99104) alternate gene name: ybbQ~similar to hypothetical proteins [Bacillus subtilis] emb CAB11968.1  (Z99105) alternate gene name: ybbQ~similar to hypothetical proteins [Bacillus subtilis] Length = 273
	11	50	09
Contig130 (59416-59787 p)	Contig130 (57941-59293 p)	Contig130 (57018-57887 p)	Contig130 (56011-56925 p)
SA-1221.1	SA-1222.1	SA-1223.1	SA-1225.1
SeqID 359	SeqID 360	SeqID 361	SeqID 362

%), Gaps = ipeptide mobilis   tripeptide   Length =	(51%), Gaps = ynthase CobQ - 02) cobyric acid Length = 252	%), Gaps = ntein ligase	3 (81%), Gaps = RIBOSOMAL omal protein L17 - somal protein L17 bosomal protein 3104) ribosomal Length = 120	%), Gaps = enase (EC um 'ogenase	%), Gaps = poamide S- Length = 436
Identities = 144/442 (32%), Positives = 229/442 (51%), Gaps = 17/442 (3%) pir  T31440 UDP-N-acetylmuramyl tripeptide synthetase homolog murC - Heliobacillus mobilis gb AAC84012.1  (AF080002) UDP-N-acetylmuramyl tripeptide synthetase MurC [Heliobacillus mobilis] Length = 455	Identities = 89/250 (35%), Positives = 129/250 (51%), Gaps = 9/250 (3%) pir  T31439 probable cobyric acid synthase CobQ - Heliobacillus mobilis gb AAC84011.1  (AF080002) cobyric acid synthase CobQ [Heliobacillus mobilis]	Identities = 153/316 (48%), Positives = 212/316 (66%), Gaps = 3/316 (0%) dbj BAB04402.1  (AP001509) lipoate-protein ligase [Bacillus halodurans]	Identities = 95/128 (74%), Positives = 105/128 (81%), Gaps = 8/128 (6%) splP20277[RL17_BACSU 50S RIBOSOMAL PROTEIN L17 (BL15) (BL21) pit  F32307 ribosomal protein L17 - Bacillus subtilis gb AAA22218.1 (M26414) ribosomal protein L17 [Bacillus subtilis] gb AAB06827.1 (L47971) ribosomal protein L17 [Bacillus subtilis] emb CAB11920.1 (299104) ribosomal protein L17 (BL15) [Bacillus subtilis]	Identities = 229/589 (38%), Positives = 339/589 (56%), Gaps = 25/589 (4%) pir  40794 dihydrolipoamide dehydrogenase (EC 1.8.1.4) [validated] - Clostridium magnum gb AAA21748.1  (L31844) dihydrolipoamide dehydrogenase [Clostridium magnum] Length = 578	Identities = 187/462 (40%), Positives = 267/462 (57%), Gaps = 26/462 (5%) dbj BAB04497.1  (AP001509) dihydrolipoamide S-acetyltransferase [Bacillus halodurans] Length = 43
50	47	63	79	57	57
Contig130 (54591-55910 m)	Contig130 (53806-54477 m)	Contig130 (52710-53699 p)	Contig131 (1559-1945 p)	Contig130 (50855-52612 p)	Contig130 (49407-50795 p)
SA-1226.1	SA-1227.1	SA-1228.1	SA-123.1	SA-1230.1	SA-1231.1
SeqID 363	SeqID 364	SeqID 365	SeqID 366	SeqID 367	SeqID 368



SeqID 369	SA-1232.1	Contig130 (48282-49280-p)	72	Identities = 189/319 (59%), Positives = 249/319 (77%), Gaps = 1/319 (0%) dbj BAB04496.1  (AP001509) acetoin dehydrogenase (TPP-dependent) beta chain [Bacillus halodurans] Length = 327
SeqID 370	SA-1233.1	 Contig130 (48230-48595 m)	. 40	Identities = 45/97 (46%), Positives = 50/97 (51%) pir  G72548 hypothetical protein APE1675 - Aeropyrum pernix (strain K1) dbj BAA80676.1  (AP000062) 155aa long hypothetical protein [Aeropyrum pernix] Length = 155
SeqID 371	SA-1234.1	Contig130 (47239-48207 p)	64	Identities = 148/317 (46%), Positives = 214/317 (66%), Gaps = 1/317 (0%) dbj BAB04495.1  (AP001509) acetoin dehydrogenase (TPP-dependent) alpha chain [Bacillus halodurans] Length = 326
SeqID 372	SA-1236.3	Contig130 (45180-47090 p)	75	Identities = 397/641 (61%), Positives = 488/641 (75%), Gaps = 17/641 (2%) gb AAK05160.1 AE006339_3 (AE006339) ABC transporter ATP binding protein [Lactococcus lactis subsp. lactis] Length = 635
SeqID 373	SA-1238.1	Contig135 (95462-95596 p)	No Hits found	
SeqID 374	SA-1239.1	Contig135 (94858-95316 p)	No Hits found	
SeqID 375	SA-1240.1	Contig 135 (94570-94797 p)	No Hits found	
SeqID 376	SA-1241.1	Contig135 (94477-94533 p)	No Hits found	
SeqID 377	SA-1242.1	Contig135 (93779-94036 m)	No Hits found	
SeqID 378	SA-1243.1	Ccntig 135 (93763-94326 p)	No Hits found	
SeqID 379	SA-1244.1	Contig135 (93063-93815 p)	No Hits found	
SeqID 380	SA-1245.1	Contig135 (92529-93014 p)	No Hits found	
ID 381	SA-1246.1	Contig135 (92196-92423 p)	No Hits found	
SeqID 382	SA-1247.1	Contig135 (89123-91921 p)	52	Identities = 157/406 (38%), Positives = 216/406 (52%), Gaps = 28/406 (6%) gb AAB30711.1  (S70345) SpaA=endocarditis immunodominant antigen [Streptococcus sobrinus, MUCOB 263, Peptide, 1566 aa] Length = 1566

$\bar{\mathcal{S}}$	Contig135 (88346-89056 p)		Identities = 85/268 (31%), Positives = 129/268 (47%), Gaps = 35/268 (13%) pir  S45091 hypothetical protein iota - Streptococcus pyogenes plasmid pDB101 pir  S68125 hypothetical protein iota - Streptococcus pyogenes plasmid pSM19035 emb CAA47097.1  (X66468) orf iota [Streptococcus pyogenes] Length = 268
Contig135 (8832	(88324-88794 m)	30	Identities = 37/104 (35%), Positives = 48/104 (45%), Gaps = 5/104 (4%) ref[XP_007094.1  collagen, type IV, alpha 1 [Homo sapiens] Length = 1669
Contig131 (60	1 (606-1544 p)	79	Identities = 190/314 (60%), Positives = 249/314 (78%), Gaps = 2/314 (0%) splO50634 RPOA_BACHD DNA-DIRECTED RNA POLYMERASE ALPHA CHAIN (TRANSCRIPTASE ALPHA CHAIN) (RNA POLYMERASE ALPHA SUBUNIT) pir  T44410 DNA-directed RNA polymerase (EC 2.7.7.6) alpha chain rpoA [imported] - Bacillus halodurans dbj BAA24194.1  (AB010082) RNA polymerase alpha subunit [Bacillus halodurans] dbj BAA75298.1  (AB017508) rpoA homologue (identity of 85 to B. subtilis%) [Bacillus halodurans] dbj BAB03881.1  (AP001507) DNA-directed RNA polymerase alpha subunit [Bacillus halodurans]
Contig135 (86100-88331 p)	0-88331 p)	45	Identities = 185/839 (22%), Positives = 342/839 (40%), Gaps = 124/839 (14%) pir  G41662 130K surface exclusion protein Sec10 precursor - Enterococcus faecalis plasmid pCF10 gb AAA65847.1  (M64978) surface exclusion protein [Plasmid pCF10] Length = 891
Contig135 (8578	(85784-86083 p) N	No Hits found	
Contig135 (8520	(85204-85401 p)	09	Identities = 29/58 (50%), Positives = 41/58 (70%) ref[NP_049430.1  unknown [Streptococcus thermophilus bacteriophage DT1] gb AAD21918.1  (AF085222) unknown [Streptococcus thermophilus bacteriophage DT1] Length =
Contig135 (85022-85207 p)	_		



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Identities = 95/192 (49%), Positives = 128/192 (66%), Gaps = 7/192 (3%) refINP_038742.1  IFN-response element binding factor 1 [Mus musculus] sp P22560 RBF1_MOUSE IFN-RESPONSE BINDING FACTOR 1 (IREBF-1) pir  A38558 interferon response element-binding factor 1 - mouse gb AAA37884.1  (M55290) IFN-response element binding factor 1 [Mus musculus]			Identities = 28/81 (34%), Positives = 46/81 (56%), Gaps = 3/81 (3%) splP21318 YR7D_ECOLI HYPOTHETICAL 11.0 KD PROTEIN (ORFD) (RETRON EC67) pir  JQ0859 hypothetical 11K protein - Escherichia coli retron Ec67 gb AAA23395.1  (M55249) unknown [Escherichia coli] Length = 100		Identities = 244/321 (76%), Positives = 286/321 (89%), Gaps = 2/321 (0%) pir  C70015 probable GMP reductase (EC 1.6.6.8) yumD - Bacillus subtilis emb CAB07955.1 (293939) unknown [Bacillus subtilis] emb CAB15203.1 (299120) similar to GMP reductase [Bacillus subtilis] Length = 326	Identities = 209/376 (55%), Positives = 286/376 (75%), Gaps = 3/376 (0%) gb AAK06013.1 AE006422_2 (AE006422) Na+/H+ antiporter [Lactococcus lactis subsp. lactis]	Identities = 86/319 (26%), Positives = 147/319 (45%), Gaps = 34/319 (10%) splP31847 YPUA_BACSU HYPOTHETICAL 31.3 KD PROTEIN IN LYSA-PPIB INTERGENIC REGION (ORFX19) pir  JU0473 ypuA protein - Bacillus subtilis gb AAA67474.1  (L09228) ORFX19 [Bacillus subtilis] emb CAB14269.1  (299116) ypuA [Bacillus subtilis] Length = 290
98	No Hits found	No Hits found	20	No Hits found	85	. 92	45
Contig135 (83986-85020 p)	Contig135 (83829-83993 p)	Contig135 (83345-83722 p)	Contig135 (82756-83013 p)	Contig135 (82201-82572 p)	Contig135 (81104-82087 p)	Contig125 (17134-18261 m)	Contig125 (16108-17070 p)
SA-1254.1	SA-1255.1	SA-1256.1	SA-1257.1	SA-1258.1	SA-1261.1	SA-1265.2	SA-1266.1
SeqID 390	SeqID 391	SeqID 392	SeqID 393	SeqID 394	SeqID 395	SeqID 396	SeqID 397

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Identities = 76/139 (54%), Positives = 92/139 (65%), Gaps = 12/139 (8%) splP45871 YWKD_BACSU HYPOTHETICAL 14.8 KD PROTEIN IN TDK-PRFA INTERGENIC REGION pir  S55436 conserved hypothetical protein ywkD - Bacillus subtilis emb CAA89883.1  (Z49782) unknown [Bacillus subtilis] emb CAB15719.1  (Z99122) similar to hypothetical proteins [Bacillus subtilis]	Identities = 65/126 (51%), Positives = 89/126 (70%), Gaps = 1/126 (0%) gb[AAA73122.1] (M77279) alpha-amylase [unidentified cloning vector]	Identities = 107/123 (86%), Positives = 115/123 (92%) sp P04969 RS11_BACSU 30S RIBOSOMAL PROTEIN S11 (BS11) pir  R3BSS1 ribosomal protein S11 - Bacillus subtilis gb AA22216.1  (M26414) ribosomal protein S11 [Bacillus subtilis] gb AA22707.1  (M13957) ribosomal protein S11 [Bacillus subtilis] gb AAB06825.1  (L47971) ribosomal protein S11 [Bacillus subtilis] emb CAB11918.1  (299104) ribosomal protein S11 (BS11) [Bacillus subtilis]	Identities = 712/819 (86%), Positives = 769/819 (92%) gb AAF63266.1  (AF220945) DNA gyrase A subunit [Streptococcus pyogenes] Length = 828	Identities = 290/329 (88%), Positives = 313/329 (94%), Gaps = 1/329 (0%) sp O33734 LDH_STRPN L-LACTATE   DEHYDROGENASE emb CAA04010.1  (AJ000336) L-lactate   dehydrogenase [Streptococcus pneumoniae]   Length = 328	Identities = 363/458 (79%), Positives = 408/458 (88%), Gaps = 3/458 (0%) gb AAC26485.2  (AF014458) NADH oxidase
64	35		92	06	. 88
Contig125 (15541-15954 p)	Contig125 (14782-15525 p)	Contig131 (173-556 p)	Contig125 (12316-14775 p)	Contig125 (11088-12077 m)	Contig125 (9598-10968 p)
SA-1267.1	SA-1268.1	SA-127.1	SA-1271.1	SA-1273.1	SA-1274.1
SeqID 398	SeqID 399	SeqID 400	SeqID 401	SeqID 402	SeqID 403

SeqID 404	SA-1275.1	Contig125 (8436-9392 p)	22	Identities = 172/318 (54%), Positives = 234/318 (73%) pir  F70009 conserved hypothetical protein yufQ - Bacillus subtilis emb CAB07939 1  (293937) unknown [Bacillus subtilis] emb CAB15146 1  (299120) similar to hypothetical proteins [Bacillus subtilis]
SeqID 405	SA-1278.1	Contig125 (7373-8434 p)	. 09	Identities = 154/349 (44%), Positives = 220/349 (62%), Gaps = 6/349 (1%) pir[IE70009 conserved hypothetical protein yufP - Bacillus subtilis emb[CAB07938.1] (Z93937) unknown [Bacillus subtilis] emb[CAB15145.1] (Z99120) similar to hypothetical proteins [Bacillus subtilis] Length = 348
SeqID 406	SA-1279.1	Contig 125 (5845-7380 p)	76	Identities = 311/497 (62%), Positives = 396/497 (79%), Gaps = 1/497 (0%) pir[ D70009 probable ABC transporter yufO - Bacillus subtilis emb CAB07937.1 (293937) unknown [Bacillus subtilis] emb CAB15144.1 (299120) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis]
SeqID 407	SA-128.1	Contig131 (3-155 p)	98	Identities = 41/50 (82%), Positives = 44/50 (88%) sp P15757 RS13_BACST 30S RIBOSOMAL PROTEIN S13 pir  R3BS3F ribosomal protein S13 - Bacillus stearothermophilus Length = 119
SeqID 408	SA-1280.1	Contig125 (4651-5700 p)	19	Identities = 164/337 (48%), Positives = 225/337 (66%), Gaps = 10/337 (2%) splO05252 YUFN_BACSU HYPOTHETICAL LIPOPROTEIN YUFN. PRECURSOR pir  C70009 ABC transporter (lipoprotein) homolog yufN - Bacillus subtilis emb CAB07936.1  (Z93937) unknown [Bacillus subtilis] emb CAB15143.1  (Z99120) similar to ABC transporter (lipoprotein) [Bacillus subtilis]
SeqID 409	SA-1281.1	Contig125 (4197-4586 p)	52	identities = 66/114 (57%), Positives = 81/114 (70%) emb CAB51906.1  (AJ237978) cytidine deaminase [Bacillus psychrophilus] Length = 136

SeqID 410	SA-1282.1		, , , ,	identities = 90/201 (44%), Positives = 144/201 (70%), Gaps = 5/201 (2%) splP37872 YBXB_BACSU HYPOTHETICAL 22.5 KDA PROTEIN IN RPLL-RPOB INTERGENIC REGION (P23) (ORF23) pir  F69751 conserved hypothetical protein ybxB - Bacillus subtilis gb AAB00971.1  (L24376) hypothetical protein [Bacillus subtilis] emb CAB11882.1  (Z99104) alternate gene name: ybaA~similar to hypothetical proteins [Bacillus subtilis] Length = 201
SeqID 411	SA-1283.1	Contig125 (2255-3175 m)	65	Identities = 140/307 (45%), Positives = 203/307 (65%), Gaps = 5/307 (1%) splQ9K8X7 COAA_BACHD PANTOTHENATE KINASE (PANTOTHENIC ACID KINASE) dbj BAB06594.1  (AP001516) pantothenate kinase [Bacillus halodurans] Length = 316
SeqID 412	SA-1284.1	Contig125 (1953-2195 m)	69	Identities = 47/86 (54%), Positives = 59/86 (67%), Gaps = 7/86 (8%) dbj BAB05058.1  (AP001511) ribosomal protein S20 (BS20) [Bacillus halodurans]
SeqID 413	SA-1285.1	Contig125 (1035-1868 p)	99	tdentities = 170/269 (63%), Positives = 203/269 (75%), Gaps = 2/269 (0%) gb AAC35851.1  (AF086736) amino acid-binding protein Abp [Streptococcus uberis]
SeqID 414	SA-1286.1	Contig125 (392-1021 p)	63	Identities = 94/210 (44%), Positives = 145/210 (68%), Gaps = 3/210 (1%) gb AAB49429.1  (U73111) glutamine transport ATP-binding protein GLNQ [Salmonella typhimurium]  Length = 240
SeqID 415	SA-1287.1	Contig125 (196-381 p)	64	Identities = 54/125 (43%), Positives = 82/125 (65%), Gaps = 1/125 (0%) dbj BAA98402.1  (AP002545) ABC amino acid transporter permease [Chlamydophila pneumoniae] Length = 217
SeqID 416	SA-1288.3	Contig139 (155214-156659 m)	21	Identities = 60/268 (22%), Positives = 107/268 (39%), Gaps = 42/268 (15%) gb AAG44891.1 AF286595_1 (AF286595) flavin-containing monooxygenase FMO3 [Rattus norvegicus] Length = 531

SeqiD 417	SA-1289.1	Contig139 (154486-155157 p)	99	Identities = 116/213 (54%), Positives = 152/213 (70%) gb AAD47593.1 AF140784_2 (AF140784) Vexp2 [Streptococcus pneumoniae] Length = 215
SeqID 418	SA-1290.1	Contig139 (154080-154274 p)	No Hits found	
SeqID 419	SA-1291.1	Contig139 (153049-154071 p)	45	Identities = 100/348 (28%), Positives = 158/348 (44%), Gaps = 17/348 (4%) gblAAD47594.1 AF140784_3 (AF140784) Vexp3 [Streptococcus pneumoniae]
SeqID 420	SA-1292.1	Contig139 (152386-153039 p)	42	Identities = 61/210 (29%), Positives = 105/210 (49%), Gaps = 18/210 (8%) pir  F69762 transporter homolog ycll - Bacillus subtilis dbj  BAA09006.1  (D50453) homologue of hypothetical protein in a rapamycin synthesis gene cluster of Streptomyces hygroscopicus [Bacillus subtilis] emb  CAB12182.1  (Z99106) similar to transporter [Bacillus subtilis]
SeqID 421	SA-1293.1	Contig139 (151707-152243 p)	83	Identities = 175/178 (98%), Positives = 176/178 (98%) gb AAF01071.1  (AF157015) CylK [Streptococcus agalactiae] Length = 178
SeqID 422	SA-1295.1	Contig139 (150488-151699 p)	94	Identities = 396/403 (98%), Positives = 400/403 (98%) gb AAF01070.1  (AF157015) CylJ [Streptococcus agalactiae] Length = 403
SeqID 423	SA-1296.1	Contig139 (148288-150483 p)	96	ldentities = 730/731 (99%), Positives = 731/731 (99%) gb AAF89495.1  (AF093787) Cyll [Streptococcus agalactiae] Length = 731
SeqID 424	SA-1297.1	Contig139 (147338-148291 p)	66	Identities = 317/317 (100%), Positives = 317/317 (100%) gb AAF89494.1  (AF093787) CylF [Streptococcus agalactiae] Length = 317
SeqID 425	SA-1298.1	Contig139 (145338-147341 p)	66	Identities = 666/667 (99%), Positives = 667/667 (99%)   gb AAD32040.1 AF093787_8 (AF093787) CylE [Streptococcus   agalactiae]
SeqID 426	SA-1299.1	Contig139 (144463-145341 p)	46	Identities = 291/292 (99%), Positives = 292/292 (99%) gb AAD32039.1 AF093787_7 (AF093787) ABC transporter homolog CylB [Streptococcus agalactiae]
SeqID 427	SA-13.1	Contig137 (32737-32958 p)	No Hits found	

SeqID 428	SA-130.1	Contig 138 (96433-96687 p)	No Hits found	
SeqID 429	SA-1300.1	Contig139 (143541-144470 p)		Identities = 308/309 (99%). Positives = 308/309 (99%) gb AAD32038 1 AF093787_6 (AF093787) ABC transporter homolog CylA [Streptococcus agalactiae] Length = 309
SeqID 430	SA-1301.1	Contig139 (143075-143551 p)	75	Identities = 120/120 (100%). Positives = 120/120 (100%) gb[AAD32037 1 AF093787_5 (AF093787) CylZ [Streptococcus agalactiae] Length = 131
SeqID 431	SA-1302.1	Contig139 (142786-143091 p)	86	7%), F 4 (A us 101
SeqID 432	SA-1303.1	Contig139 (142071-142793 p)	94	
SeqID 433	SA-1304.2	Contig139 (141226-142074 p)	66	Identities = 282/282 (100%), Positives = 282/282 (100%) gbjAAD32034.1jAF093787_2 (AF093787) CylD [Streptococcus agalactiae]
SeqID 434	SA-1305.2	Contig139 (140921-141226 p)	69	Identities = 101/101 (100%), Positives = 101/101 (100%) gb AAD32033.1 AF093787_1 (AF093787) CylX [Streptococcus agalactiae]
SeqID 435	SA-1308.2	Contig128 (27763-28422 p)	40	Identities = 54/181 (29%), Positives = 95/181 (51%), Gaps = 1/181 (0%) emb CAA65740.1  (X97014) PrfA [Listeria seeligeri] Length = 237
SeqID 436	SA-1309.1	Contig128 (28446-30731 p)	65	Identitles = 386/767 (50%), Positives = 502/767 (65%), Gaps = 21/767 (2%) splP22093 PEPX_LACLC XAA-PRO DIPEPTIDYL-PEPTIDASE (X-PRO DIPEPTIDYL-PEPTIDASE) (X-PROLYL-DIPEPTIDYL AMINOPEPTIDASE) (X-PDAP) pir  A43747 X-Pro dipeptidyl-peptidase (EC 3.4.14.11) - Lactococcus lactis subsp. cremoris gb AAA25232.1  (M58315) dipeptidyl peptidase IV [Lactococcus lactis] Length = 763
SeqID 437	SA-1310.1	Contig128 (30735-31094 p)	No Hits found	



1 0 T G	n bead log	a · sa c · · · · · · · · · · · · · · · · · ·
Identities = 101/318 (31%), Positives = 184/318 (57%), Gaps = 5/318 (1%) splP31114 HEP2_BACSU HEPTAPRENYL DIPHOSPHATE SYNTHASE COMPONENT II (HEPPP SYNTHASE COMPONENT II (HEPPP SYNTHASE SUBUNIT 2) (SPORE GERMINATION PROTEIN C3) pir  E69630 heptaprenyl diphosphate synthase component II gerCC - Bacillus subtilis gb AAA20856.1  (M80245) GerC3 [Bacillus subtilis] emb CAB14190.1  (Z99115) heptaprenyl diphosphate synthase component II [Bacillus subtilis]	Identities = 264/577 (45%), Positives = 391/577 (67%), Gaps = 14/577 (2%) splP94367 CYDD_BACSU TRANSPORT ATP-BINDING PROTEIN CYDD_pir ID69611 ABC transporter required for expression of cytochrome bd (ATP-) cydD - Bacillus subtilis dbj BAA11730.1  (D83026) homologous to many ATP-binding transport proteins; hypothetical [Bacillus subtilis] emb CAB15899.1  (299123) ABC membrane transporter (ATP-binding protein) [Bacillus subtilis] Length = 575	Identities = 278/569 (48%); Positives = 399/569 (69%), Gaps = 6/569 (1%) splP94366 CYDC_BACSU TRANSPORT ATP-BINDING PROTEIN CYDC pir  C69611 ABC transporter required for expression of cytochrome bd (ATP-) cydC - Bacillus subtilis dbj BAA11729.1  (D83026) homologous to many ATP-binding transport proteins including SwissProt:CYDD_ECOLI; hypothetical [Bacillus subtilis] emb CAB15900.1  (Z99123) ABC membrane transporter (ATP-binding protein) [Bacillus subtilis]
IS (57% PTAPR VT II (H SERMIN Sphate gb AA/4190.1	7 (67% ansport cydD cydD cydD us to m [Bacillu ransport Lengton Lengton ransport cydD cydD cydD cydD cydD cydD cydD cydD	7669 (69% RANSPOR ; transport cydC gous to m including iillus iillus ie transpoi
tives = 184/31 2_BACSU HE E COMPONE 2) (SPORE G taprenyl dipho subtilis s] emb[CAB1 thase compon	: 391/57 SU TRA ABC tr ABC tr ) mologo mologo thetical thetical	399/56 U TRAI U TRAI ABC tr ABC tr mologo mologo inc
sitives = 22_BAC SE CON T 2) (SF ptapren tis] em tihase c	sitives = 30 De9611 At De9611 At d (ATP-) D26) homo hypothe SC membri	sitives = 36 BACSU 7 C69611 AE d (ATP-) 226) homo teins thetical [B thetical [B thetical [B thetical [B
entities = 101/318 (31%), Positives = 184/318 (57%), Gaps 5/318 (1%) splP31114 HEP2_BACSU HEPTAPRENYL DIPHOSPHATE SYNTHASE COMPONENT II (HEPPP SYNTHASE COMPONENT II (HEPPP SYNTHASE SUBUNIT 2) (SPORE GERMINATION ROTEIN C3) pir  E69630 heptaprenyl diphosphate synthamponent II gerCC - Bacillus subtilis] emb CAB14190.1  (2991 heptaprenyl diphosphate synthase component II [Bacillus subtilis]	%), Pos s7[CYDI 3D pir[] nrome b 11 (D83 ns; ns; lus	278/569 (48%); Positives %) splP94366 CYDC_BACROTEIN CYDC pirl C696 sion of cytochrome bd (Alg) BAA11729.1  (D83026) binding transport proteins CYDD_ECOLI; hypothetic 5900.1  (299123) ABC me rotein) [Bacillus
318 (31%) ATE SYN SUP DIT SYN SUP DIT SYN SUP	lentities = 264/577 (45%) 14/577 (2%) splP94367 (40) 14/577 (2%) splP94367 (40) 14/577 (2%) splP94367 (40) splP94367 (45%) spl	tentities = 278/569 (48%) 6/569 (1%) splP94366 C 4DING PROTEIN CYDC or expression of cytochrolubtilis dbj BAA11729.1  binding transporwissProt:CYDD_ECOLI; mb CAB15900.1  (29912) binding protein) [Bacillus
entities = 101/ 5/318 (1%); DIPHOSPH, SYNTHASE ROTEIN C3) Imponent II ge 80245) GerC3 heptaprenyl c	s = 264/ (2%) st PROTE ession o dbjlBAA ranspor ranspor ranspor	= 278/ 1%) sp PROTE sssion o dbj BAA dbj BaA d
Identities = 101/318 (31%), Pos 5/318 (1%) splP31114 HEP DIPHOSPHATE SYNTHAS SYNTHASE SUBUNI PROTEIN C3) pir  E69630 hej component II gerCC - Bacillus M80245) GerC3 [Bacillus subti heptaprenyl diphosphate syr	Identities = 264/577 (45%), Positives = 391, 14/577 (2%) splP94367 CYDD_BACSU TINDING PROTEIN CYDD pir  D69611 ABC for expression of cytochrome bd (ATP-) subtilis dbj  BA411730.1  (D83026) homolo binding transport proteins; hypothetic emb  CAB15899.1  (299123) ABC membran binding protein) [Bacillus subtilis]	Identities = 278/569 (48%); Positives = 399/569 6/569 (1%) splP94366 CYDC_BACSU TRANSINDING PROTEIN CYDC_pir  C69611 ABC transfor expression of cytochrome bd (ATP-) subtilis dbj  BAA11729.1  (D83026) homologous binding transport proteins inclusingsProt:CYDD_ECOLI; hypothetical [Bacillus emb  CAB15900.1  (299123) ABC membrane trabinding protein) [Bacillus
	_ <u> </u>	<u> </u>
99	99	29
20 p)	33 m)	£ E
40-321:	05-3396	46-3566
28 (311	28 (3220	78 (338 <sub>4</sub>
Contig128 (31140-32120 p)	Contig128 (32205-33953 m)	Contig128 (33946-35664 m)
SA-1311.1	SA-1312.1	SA-1313.1
SeqID 438	SeqID 439	SeqID 440
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Identities = 158/331 (47%), Positives = 223/331 (66%), Gaps = 1/331 (0%) splP94365 CYDB_BACSU CYTOCHROME D UBIQUINOL OXIDASE SUBUNIT II pir  B69611 cytochrome d ubiquinol oxidase (EC 1.10.3) chain Il cydB - Bacillus subtilis emb CAB15901.1  (D83026) homologous to cytochrome d ubiquinol oxidase subunit II; hypothetical [Bacillus subtilis] emb CAB15901.1  (Z99123) cytochrome bd ubiquinol oxidase (subunit II) [Bacillus subtilis] Length = 338	Identities = 246/470 (52%), Positives = 319/470 (67%), Gaps = 12/470 (2%) splP94364 CYDA_BACSU CYTOCHROME DUBIQUINOL OXIDASE SUBUNIT I pir  A69611 cytochrome bdubiquinol oxidase (subunit I) cydA - Bacillus subtilis dbj  BAA11727.1  (D83026) homologous to cytochrome dubiquinol oxidase subunit I; hypothetical [Bacillus subtilis] emb  CAB15902.1  (Z99123) cytochrome bd ubiquinol oxidase (subunit I) [Bacillus subtilis] Length = 468	Identities = 178/403 (44%), Positives = 249/403 (61%), Gaps = 7/403 (1%) pir  A70015 probable NADH dehydrogenase (EC 1.6.99.3) yumB - Bacillus subtilis emb CAB07953.1  (293939) unknown [Bacillus subtilis] emb CAB15200.1  (299120) similar to NADH dehydrogenase [Bacillus subtilis] Length = 406	
SA-1314.1	SA-1316.1	SA-1318.1 C	
SeqID 441	SeqID 442	SeqID 443	



				= 220 / V4Th / O00/95 = positional / V3E/ O00/17 = positional
Ø	SA-1319.2	Contig128 (39435-40334 m)	4	to quinone  15/290 (5%) splP39582 MENA_BACSU PROBABLE 1,4-  15/290 (5%) splP39582 MENA_BACSU PROBABLE 1,4-  DIHYDROXY-2-NAPHTHOATE OCTAPRENYLTRANSFERASE  (DHNA-OCTAPRENYLTRANSFERASE) pir  S39661  menaquinone biosynthesis protein homolog ywaB - Bacillus  subtilis emb CAA51562.1  (X73124) ipa-6d [Bacillus subtilis]  emb CAB15875.1  (Z99123) alternate gene name: ipa-6d~similar  to quinone  biosynthesis [Bacillus subtilis]  Length =  311
•	SA-132.2	Contig138 (94727-95020 m)	42	Identities = 35/72 (48%), Positives = 42/72 (57%) pirglS59084 hypothetical protein 29.1 - red alga (Chondrus crispus) mitochondrion emb[CAA87600.1] (Z47547) unique orf [Chondrus crispus] Length = 79
יטן	SA-1320.1	Contig100 (14165-14662 p)	No Hits found	
တ	SA-1322.1	Contig 100 (13666-14115 p)	No Hits found	
07	SA-1323.1	Contig100 (13349-13642 p)	No Hits found	
0,	SA-1324.1	Contig100 (12954-13271 p)	29	Identities = 70/96 (72%), Positives = 83/96 (85%) gb AAB52379.1  (U31811) immunogenic secreted protein precursor [Streptococcus pyogenes] Length = 534
<i>o</i>	SA-1325.1	Contig100 (12107-13018 p)	09	Identities = 142/298 (47%), Positives = 185/298 (61%), Gaps = 11/298 (3%) gb AAB52379.1  (U31811) immunogenic secreted protein precursor [Streptococcus pyogenes] Length = 534
0)	SA-1326.1	Contig100 (11805-12203 p)	32	Identities = 35/112 (31%), Positives = 49/112 (43%), Gaps = 12/112 (10%) pir  T07945 probable arabinogalactan protein (clone Sta 39-3) - rape gb AAC37509.1  (L47351) arabinogalactan protein [Brassica napus] Length = 136
တျ	SA-1327.1	Contig100 (11394-11648 p)	57	Identities = 32/76 (42%), Positives = 54/76 (70%) dbj BAB04699.1  (AP001510) unknown conserved protein [Bacillus halodurans] Length = 102

SeqID 453	SA-1328.1	Contig100 (10783-11373 p)	55	Identities = 72/259 (27%), Positives = 112/259 (42%), Gaps = 64/259 (24%) pir  C82882 ABC Transporter UU510 [imported] - Ureaplasma urealyticum gb AAF30922.1 AE002149] ABC Transporter [Ureaplasma urealyticum]
SeqID 454	SA-1329.1	Contig100 (10314-10769 p)	No Hits found	
SeqID 455	SA-133.1		63	Identities = 62/104 (59%), Positives = 68/104 (64%) pir  F81516 hypothetical protein CP0987 [imported] - Chlamydophila pneumoniae (strain AR39) Length = 216
SeqID 456	SA-1330.1	Contig100 (9421-10335 p)	34	Identities = 59/254 (23%), Positives = 108/254 (42%), Gaps = 14/254 (5%) emb CAC16670.1  (AJ302698) hypothetical protein [Staphylococcus haemolyticus] Length = 373
SeqID 457	SA-1331.1	Contig100 (9062-9418 p)	No Hits found	
SeqID 458	SA-1332.1	Contig100 (8700-9050 p)	No Hits found	
SeqID 459	SA-1333.2	Contig100 (8666-8776 m)	No Hits found	
SeqID 460	SA-1334.1	Contig100 (4757-8686 p)	20	Identities = 452/1058 (42%), Positives = 664/1058 (62%), Gaps = 39/1058 (3%) pir  C70013 conserved hypothetical protein yukA - Bacillus subtilis emb CAB15175.1  (299120) alternate gene name: yueA~similar to hypothetical proteins [Bacillus subtilis] . Length = 1207
SeqID 461	SA-1335.1	Contig100 (4279-4782 p)	No Hits found	
SeqID 462	SA-1336.1	Contig100 (2998-4272 p)	37	Identities = 83/323 (25%), Positives = 163/323 (49%), Gaps = 1/323 (0%) dbj BAB04693.1  (AP001510) unknown conserved protein [Bacillus halodurans]
SeqID 463	SA-1337.1	Contig100 (2756-2998 p)	No Hits found	
SeqID 464	SA-1338.1	Contig100 (2344-2772 p)	No Hits found	
SeqID 465	SA-1339.1	Contig100 (2-2290 p)	27	Identities = 101/501 (20%), Positives = 213/501 (42%), Gaps = 54/501 (10%) pir  PC6003 surface membrane protein Imp4 - Mycoplasma hominis (fragment) Length = 624
SeqID 466	SA-134.1	Contig138 (94635-94964 m)	No Hits found	



E 5- UCTASE) ergenic 14003) 97) 5-keto- Length =	ps = rase 171	Gaps = nase - ɔ-3- Length =	aps = (M0066)	aps = protein	aps = orted] - rotein
Identities = 116/260 (44%), Positives = 165/260 (62%), Gaps = 6/260 (2%) splP39345 IDNO_ECOLI GLUCONATE 5-DEHYDROGENASE (5-KETO-D-GLUCONATE 5-REDUCTASE) pir  S56492 hypothetical 27.6K protein (pepa-gntv intergenic region) - Escherichia coli gb AAA97163.1  (U14003) ORF_f254 [Escherichia coli] gb AAC77223.1  (AE000497) 5-keto-D-gluconate 5-reductase [Escherichia coli K12] Length = 254	Identities = 47/189 (24%), Positives = 81/189 (41%), Gaps = 20/189 (10%) pir  D43258 galactose-6-phosphate isomerase subunit LacB - Streptococcus mutans Length = 171	Identities = 94/329 (28%), Positives = 169/329 (50%), Gaps 7/329 (2%) pir  G72422 2-keto-3-deoxygluconate kinase - Thermotoga maritima (strain MSB8) gb AAD35161.1 AE001693_7 (AE001693) 2-keto-3-deoxygluconate kinase [Thermotoga maritima] Length 339	Identities = 78/192 (40%), Positives = 118/192 (60%), Gaps = 6/192 (3%) pir F72422 KHG-KDPG bifunctional aldolase TM0066 [similarity] - Thermotoga maritima (strain MSB8) gb AAD35160.1 AE001693_6 (AE001693) 2-dehydro-3-deoxyphosphogluconate aldolase/4-hydroxy-2-oxoglutarate aldolase [Thermotoga maritima] Length = 205	Identities = 112/529 (21%), Positives = 218/529 (41%), Gaps = 67/529 (12%) emb CAB62846.2  (AL035475) hypothetical protein [Plasmodium falciparum]	Identities = 115/203 (56%), Positives = 143/203 (69%), Gaps = 7/203 (3%) pirl A83323 hypothetical protein PA2575 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG05963.1 AE004686_1 (AE004686) hypothetical protein [Pseudomonas aeruginosa] Length = 200
<u> </u>		- 0	0 ×0	)   67	9
28	38	48	51	35	29
.268 m)	439 m)	.775 m)	756 m)	1410 m)	1386 p)
Contig120 (16456-17268 m)	Contig120 (15801-16439 m)	Contig120 (14768-15775 m)	Contig120 (14118-14756 m)	- Contig120 (11557-13410 m)	Contig120 (10772-11386 p)
SA-1340.2	SA-1341.1	SA-1342.1	SA-1343.1	SA-1344.2	SA-1345.1
SeqID 467	SeqID 468	SeqID 469	SeqID 470	SeqID 471	SeqID 472

Identities = 49/124 (39%), Positives = 73/124 (58%) pir  A69860 transcription regulator MarR family homolog ykoM - Bacillus subtilis emb CA405611.1  (AJ002571) YkoM [Bacillus subtilis] emb CA813191.1  (Z99110) similar to transcriptional regulator (MarR family) {Bacillus subtilis} emb CA813207.1  (Z99111) similar to transcriptional regulator (MarR family) {Bacillus subtilis}	Identities = 1238/1468 (84%), Positives = 1346/1468 (91%), Gaps = 3/1468 (0%) gb AAF98345.1  (AF280761) DNA polymerase III alpha chain [Streptococcus pyogenes] Length = 1465	Identities = 101/201 (50%), Positives = 122/201 (60%), Gaps   9/201 (4%) emb CAB94815.1  (AJ245582) peptidoglycan hydrolase [Streptococcus thermophilus] Length = 218	No Hits found	Identities = 300/608 (49%), Positives = 410/608 (67%), Gaps = 52/608 (8%) sp O31755 SYP_BACSU PROLYL-TRNA SYNTHETASE (PROLINETRNA LIGASE) (PRORS) pir  G69682 prolinetRNA ligase (EC 6.1.1.15) proS - Bacillus subtilis emb CAB13530.1  (Z99112) prolyl-tRNA synthetase [Bacillus subtilis]	Identities = 229/425 (53%), Positives = 298/425 (69%), Gaps = 9/425 (2%) gb AAD47948.1 AF152237_1 (AF152237) Eep [Enterococcus faecalis] Length = 422	Identities = 116/266 (43%), Positives = 172/266 (64%), Gaps = 6/266 (2%) dbj BAB06141.1  (AP001515) phosphatidate cytidylyltransferase [Bacillus halodurans]
Contig120 (10214-10639 p)	Contig120 (5681-10087 p)	Contig120 (4974-5558 p)	Contig 138 (93460-93819 p) No H	Contig120 (2996-4849 p)	Contig120 (1645-2904 p)	Contig120 (820-1614 p)
SA-1347.1	SA-1348.1	SA-1349.1	SA-135.1	SA-1350.1	SA-1351.1	SA-1352.1
SeqiD 473	SeqID 474	SeqID 475	SeqID 476	SeqID 477	SeqID 478	SeqID 479



s = (iiis		II GO	"Ö" ∢∢ TΩ"	aps = Length		s in cia
identities = 134/235 (57%), Positives = 170/235 (72%), Gaps = 4/235 (1%) spj031751jUPPS_BACSU UNDECAPRENYL PYROPHOSPHATE SYNTHETASE (UPP SYNTHETASE) (DI-TRANS-POLY-CIS-DECAPRENYLCISTRANSFERASE) (UNDECAPRENYL DIPHOSPHATE SYNTHASE) (UDS) pirl A69881 conserved hypothetical protein yluA - Bacillus subtilis emb CAB13526.1  (299112) similar to hypothetical proteins [Bacillus subtilis]		dentities = 289/478 (60%), Positives = 368/478 (76%), Gaps = 2/478 (0%) gb AAA69226.1  (U29579) 6-phospho-beta-glucosidase [Escherichia coli]	Identities = 43/177 (24%), Positives = 84/177 (47%), Gaps = 9/177 (5%) spl035264 PA1B_RAT PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE IB BETA SUBUNIT (PAF ACETYLHYDROLASE 30 KDA SUBUNIT) (PAF-AH BETA SUBUNIT) (PAF-AH BETA SUBUNIT) (PAF-AH BETA SUBUNIT) (PLATELET-ACTIVATING FACTOR ACETYLHYDROLASE ALPHA 2 SUBUNIT) (PAF-AH ACETYLHYDROLASE ALPHA 2 SUBUNIT) (PAF-AH ALPHA 2) gb AAC27974.1  (AF016048) platelet-activating factor acetylhydrolase alpha 2 subunit [Rattus norvegicus] Length = 229	Identities = 103/265 (38%), Positives = 154/265 (57%), Gaps = 4/265 (1%) splP75809 YBJI_ECOLI PROTEIN YBJI Lengt = 271	Identities = 197/296 (66%), Positives = 240/296 (80%) gb AAF89977.1 AF206272_3 (AF206272) transcriptional regulator [Streptococcus mutans] Length = 301	Identities = 141/443 (31%), Positives = 241/443 (53%), Gaps = 20/443 (4%) pir  C82449 conserved hypothetical protein VCA0526 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF96429.1  (AE004383) conserved hypothetical protein [Vibrio cholerae] Length = 468
63	No Hits found	92	40	55 .	75	46
Contig120 (53-805 p)	Contig99 (17810-18058 p)	Contig99 (16154-17593 p)	Contig99 (15380-15994 p)	Contig99 (14458-15279 p)	Contig99 (12904-13836 m)	Contig99 (11315-12847 p)
SA-1353.1	SA-1354.1	SA-1355.1	SA-1356.1	SA-1357.1	SA-1358.1	SA-1359.1
SeqID 480	SeqID 481	SeqID 482	SeqID 483	SeqID 484	SeqID 485	SeqID 486

SeqID 487	SA-136.1	Contig138 (92309-93091 m)	No Hits found	
SeqID 488	SA-1361.1	 Contig99 (10178-11206 p)	90	Identities = 131/336 (38%), Positives = 188/336 (54%), Gaps = 7/336 (2%) splP23861 POTD_ECOLI SPERMIDINE/PUTRESCINE-BINDING PERIPLASMIC PROTEIN PRECURSOR (SPBP) pir  D40840 spermidine/putrescine-binding protein precursor [validated] - Escherichia coli gb AAC37041.1  (M64519) transport protein [Escherichia coli] dbj BAA35943.1  (D90747) Spermidine/putrescine transport protein D [Escherichia coli] gb AAC74207.1  (AE000212) spermidine/putrescine periplasmic transport protein [Escherichia coli] 348
SeqID 489	SA-1362.1	Contig99 (9364-10140 p)		Identities = 97/249 (38%), Positives = 159/249 (62%), Gaps = 3/249 (1%) pir[ G70179 spermidine/putrescine ABC transporter, permease protein (potC) homolog - Lyme disease spirochete gb[AAB91527.1] (AE001165) spermidine/putrescine ABC transporter, permease protein (potC) [Borrelia burgdorferi] Length = 263
SeqID 490	SA-1363.1	Contig99 (8573-9367 p)	99	Identities = 90/255 (35%), Positives = 153/255 (59%), Gaps = 11/255 (4%) splP45170 POTB_HAEIN SPERMIDINE/PUTRESCINE TRANSPORT SYSTEM PERMEASE PROTEIN POTB pir  A64118 spermidine/putrescine transport system permease potB - Haemophilus influenzae (strain Rd KW20) gb AAC22990.1  (U32813) spermidine/putrescine ABC transporter, permease protein (potB) [Haemophilus influenzae Rd] Length = 286
SeqID 491	SA-1364.1	Contig99 (7435-8589 p)	. 62	Identities = 165/345 (47%), Positives = 240/345 (68%), Gaps = 1/345 (0%) pir[ A70180 spermidine/putrescine ABC transporter, ATP-binding protein (potA) homolog - Lyme disease spirochete gb AAB91525.1  (AE001165) spermidine/putrescine ABC transporter, ATP-binding protein (potA) [Borrelia burgdorferi] Length = 347

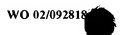


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SeqID 497	SA-1370.1	Contig99 (4121-4684 p)	87	Identities = 148/184 (80%), Positives = 168/184 (90%) sp O33723 GCH1_STRPY GTP CYCLOHYDROLASE I (GTP-CHI) emb CAA04237.1  (AJ000685) GTP cyclohydrolase [Streptococcus pyogenes] Length = 194
SeqID 498	SA-1371.1	Contig99 (2840-4102 p)	57	Identitles = 154/426 (36%), Positives = 245/426 (57%), Gaps = 17/426 (3%) sp Q05865 FOLC_BACSU   FOLYLPOLYGLUTAMATE SYNTHASE (FOLYLPOLY-GAMMA-GLUTAMATE SYNTHETASE)   polyglutamate synthetase folC - Bacillus subtilis gb AAB59021.1  (L04520) folyl-polyglutamate synthetase folC - Bacillus subtilis]   emb CAB14768.1  (Z99118) folyl-polyglutamate synthetase   [Bacillus subtilis]   Length = 430
SeqID 499	SA-1372.1	Contig99 (1951-2838 p)	63	Identities = 110/295 (37%), Positives = 188/295 (63%), Gaps = 6/295 (2%) dbj BAB07585.1  (AP001520) unknown conserved protein [Bacillus halodurans] Length = 308
SeqID 500	SA-1373.2	Contig99 (1098-1964 p)	62	Identities = 188/289 (65%), Positives = 232/289 (80%), Gaps = 1/289 (0%) splP72535 KHSE_STRPN HOMOSERINE KINASE (HK) gb AAC44297.1  (U41735) homoserine kinase homolog [Streptococcus pneumoniae] Length = 289
SeqID 501	SA-1374.2	Contig99 (2-1096 p)	71	Identities = 187/368 (50%), Positives = 261/368 (70%), Gaps = 11/368 (2%) splP52985 DHOM_LACLA HOMOSERINE DEHYDROGENASE (HDH) pir  JC6049 homoserine dehydrogenase (EC 1.1.1.3) - Lactococcus lactis emb CAA65713.1  (X96988) hom {Lactococcus lactis} Length = 428
SeqID 502	SA-1377.1	Contig134 (51276-52901 p)	59	Identities = 210/541 (38%), Positives = 326/541 (59%), Gaps = 14/541 (2%) dbj BAB04286.1  (AP001509) nickel transport system (nickel-binding protein) [Bacillus halodurans]



SeqID 503	SA-1378.1	Contig134 (52888-53832 p)	55	Identities = 121/304 (39%), Positives = 176/304 (57%) dbj BAB04287.1  (AP001509) nickel transport system (permease) [Bacillus halodurans] Length = 314
SeqID 504	SA-1379.1	Contig134 (53871-54638 p)	89	Identities = 106/255 (41%), Positives = 164/255 (63%) dbj BAB04288.1  (AP001509) nickel transport system (permease) [Bacillus halodurans] Length = 277
SeqID 505	SA-138.1	Contig138 (90969-91175 p)	No Hits found	
SeqID 506	SA-1380.1	Contig134 (54614-55414 p)	25	Identities = 85/253 (33%), Positives = 154/253 (60%), Gaps = 2/253 (0%) gb AAF73561.1 (AE002315) peptide ABC transporter, ATP-binding protein [Chlamydia muridarum]
SeqID 507	SA-1381.1	Contig134 (55401-56081 p)	55	Identities = 81/199 (40%), Positives = 131/199 (65%), Gaps = 2/199 (1%) dbj BAB05797.1  (AP001514) oligopeptide ABC transporter (ATP-binding protein) [Bacillus halodurans] Length = 244
SeqID 508	SA-1382.1	Contig134 (56202-56930 p)	76	Identities = 143/238 (60%), Positives = 193/238 (81%) sp O31749 PYRH_BACSU URIDYLATE KINASE (UK) (URIDINE MONOPHOSPHATE KINASE) (UMP KINASE) pir  F69708 uridylate kinase smbA - Bacillus subtilis emb CAB13524.1  (299112) uridylate kinase [Bacillus subtilis] Length = 240
SeqID 509	SA-1383.1	Contig134 (56946-57503 p)	78	Identities = 112/185 (60%), Positives = 149/185 (80%) dbj BAB06143.1  (AP001515) ribosome recycling factor [Bacillus halodurans] Length = 185
SeqID 510	SA-1384.1	Contig134 (57621-58475 p)	49	Identities = 107/269 (39%), Positives = 155/269 (56%), Gaps = 6/269 (2%) pir  E69840 hypothetical protein yitL - Bacillus subtilis emb CAB12943.1  (299109) yitL [Bacillus subtilis] 298
SeqID 511	SA-1385.1	Contig134 (58601-59122 p)	29	Identities = 102/175 (58%), Positives = 127/175 (72%), Gaps = 2/175 (1%) dbj BAB05167.1  (AP001512) peptide methionine sulfoxide reductase [Bacillus halodurans] Length = 179

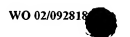


SeqID 512	SA-1386.1	Contig134 (59143-59334 p)	90	Identities = 24/66 (36%). Positives = 42/66 (63%) pirt  A69931     hypothetical protein yozE - Bacillus subtilis emb CAB13859.1      (299114) yozE [Bacillus subtilis]   Length = 74
SeqID 513	SA-1388.1	Contig134 (59492-61264 p)	68	Identities = 472/590 (80%). Positives = 535/590 (90%) gb AAA51870 1  (U09352) 67 kDa Myosin-crossreactive streptococcal antigen [Streptococcus pyogenes] Length = 590
SeqID 514	SA-1389.1	Contig134 (61388-62374 p)	99	Identities = 191/305 (62%), Positives = 241/305 (78%), Gaps = 1/305 (0%) splP46343 PHOL_BACSU PHOH-LIKE PROTEIN pir  E69676 phosphate starvation-induced protein phoH - Bacillus subtilis dbj BAA12477.1  (D84432) YqfE [Bacillus subtilis] emb CAB14476.1  (Z99117) phosphate starvation-induced protein [Bacillus subtilis] Length = 319
SeqID 515	SA-1390.2	Contig134 (62465-63268 m)	25	Identities = 42/161 (26%), Positives = 72/161 (44%), Gaps = 19/161 (11%) gb AAC95491.1  (U73025) unknown [Staphylococcus aureus] gb AAG42230.1 AF299292 (AF299292) ORFX [Staphylococcus intermedius] Length = 289
SeqID 516	SA-1391.2	Contig133 (46310-47965 m)	56	Identities = 176/589 (29%), Positives = 315/589 (52%), Gaps = 47/589 (7%) dbj BAB06875.1  (AP001517) two-component sensor histidine kinase involved in phosphate regulation [Bacillus halodurans] Length = 589
SeqID 517	SA-1392.1	Contig133 (47958-48635 m)	59	Identities = 98/224 (43%), Positives = 138/224 (60%), Gaps = 2/224 (0%) splP45606 PHOB_SHIDY PHOSPHATE REGULON TRANSCRIPTIONAL REGULATORY PROTEIN PHOB pir  A44753 phoB protein - Shigella dysenteriae gb AAA26535.1  (M31793) phosphate regulatory protein phoB [Shigella dysenteriae] Length = 229



Identities = 66/213 (30%), Positives = 119/213 (54%), Gaps = 4/213 (1%) sp[Q51547]PHOU_PSEAE PHOSPHATE TRANSPORT SYSTEM PROTEIN PHOU pir  S65576 negative regulator PhoU - Pseudomonas aeruginosa pir  S68596 negative regulator PhoU - Pseudomonas aeruginosa pir  H82975 phosphate uptake regulatory protein PhoU PA5365 [imported] - Pseudomonas aeruginosa (strain PAO1) dbj  BAA08138.1  (D45195) a negative regulator of pho regulon [Pseudomonas aeruginosa] gb  AAG08750.1 AE004948_6 (AE004948) phosphate uptake regulatory protein PhoU [Pseudomonas aeruginosa] Length = 242	Identities = 166/245 (67%), Positives = 211/245 (85%), Gaps = 1/245 (0%) gb AAD22041.1  (AF118229) ATP-binding cassette protein PstB [Streptococcus pneumoniae] Length = 250	Identities = 135/263 (51%), Positives = 203/263 (76%) gb AAD22040.1  (AF118229) transmembrane protein PstA [Streptococcus pneumoniae] Length = 271	Identities = 162/266 (60%), Positives = 212/266 (78%), Gaps = 3/266 (1%) gb/AAD22039.1 (AF118229) transmembrane protein PstC [Streptococcus pneumoniae] Length = 271		Identities = 230/230 (100%), Positives = 230/230 (100%) pir  A61607 probable hemolysin precursor - Streptococcus agalactiae (strain 74-360) Length = 230	Identities = 31/118 (26%), Positives = 51/118 (42%), Gaps = 3/118 (2%) pir  S67490 single-stranded DNA-binding protein - Eubacterium sp gb AAA79866.1  (U12515) single-stranded DNA binding protein [uncultured eubacterium] prf  2108276A
Identities = 66/213 (30%),  4/213 (1%) splQ51547  TRANSPORT SYSTEM PR regulator PhoU - Pseudomol regulator PhoU - Pseudo phosphate uptake regulator Pseudomonas aeruginos (D45195) a negative regula aeruginosa] gb AAG08 phosphate uptake regula	Identities = 166/245 (67%), 1/245 (0%) gb AAD22041.1 protein PstB [Streptococcus		Identities = 162/266 (60%), Positives = 3/266 (1%) gb AAD22039.1   (AF118229 PstC [Streptococcus pneumoniae]		Identities = 230/230 pir  A61607 probable agalactiae (strain	Identities = 31/118 (26%), Positives = 51/3/118 (2%) pir  S67490 single-stranded DN Eubacterium sp gb AAA79866.1  (U12515)
15	82	99	74	No Hits found	77	32
Contig133 (48635-49291 m)	Contig133 (49288-50037 m)	Contig133 (50030-50908 m)	Contig133 (50910-51755 m)	Contig133 (52368-52664 p)	Contig133 (51770-52651 m)	Contig137 (32042-32473 p)
SA-1393.1	SA-1394.1	SA-1395.1	SA-1396.1	SA-1398.2	SA-1399.1	SA-14.1
SeqID 518	SeqID 519	SeqID 520	SeqID 521	SeqID 522	SeqID 523	SeqID 524

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SeqID 525	SA-140.1	<u> </u>	47	Identities = 86/278 (30%), Positives = 137/278 (48%), Gaps = 36/278 (12%) dbj BAA07788.1  (D43692) rep protein [Brevibacillus borstelensis] Length = 281
SeqID 526	SA-1400.1	Contig133 (52852-53439 m)	No Hits found	
SeqID 527	SA-1401.1	Contig133 (53436-54176 m)	28	Identities = 119/250 (47%), Positives = 149/250 (59%), Gaps = 9/250 (3%) dbj BAB05069.1  (AP001511) unknown conserved protein [Bacilius halodurans] Length = 250
SeqID 528	SA-1403.1	Contig133 (54176-55129 m)	69	Identities = 138/309 (44%), Positives = 193/309 (61%), Gaps = 5/309 (1%) pir] T43740 probable ribosomal protein L11 methyltransferase (EC 2.1.1) [imported] - Listeria monocytogenes dbj BAA82791.1  (AB023064) orf35 [Listeria monocytogenes] Length = 314
SeqID 529	SA-1404.1	Contig133 (55126-55434 m)	No Hits found	
SeqID 530	SA-1405.1	Contig133 (55708-56424 p)	25	Identities = 87/246 (35%), Positives = 140/246 (56%), Gaps = 13/246 (5%) pir  C69661 transcription activator of multidrug-efflux transporter genes mta - Bacillus subtilis emb CAA69863.1  (Y08559) Unknown [Bacillus subtilis] emb CAB15677.1  (Z99122) transcriptional regulator [Bacillus subtilis]
SeqID 531	SA-1406.1	Contig133 (56463-56933 m)	36	Identities = 38/136 (27%), Positives = 60/136 (43%), Gaps = 3/136 (2%) splP54441 YRKN_BACSU HYPOTHETICAL 21.3 KD PROTEIN IN BLTR-SPOIIIC INTERGENIC REGION pir  D69977 hypothetical protein yrkN - Bacillus subtilis dbj BAA12369.1  (D84432) YrkN [Bacillus subtilis] emb CAB14586.1  (299117) yrkN [Bacillus subtilis]
SeqID 532	SA-1407.1	Contig133 (56905-57363 m)	52	Identities = 57/131 (43%), Positives = 82/131 (62%) pir  F83247 hypothetical protein PA3180 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG06568.1 AE004742_4 (AE004742) hypothetical protein [Pseudomonas aeruginosa] Length = 145
SeqID 533	SA-1408.1	134	No Hits found	
eqID 534	SA-141.1	Contig 138 (89680-89994 p)	No Hits found	



SeqID 535	SA-1410.1	Contig133 (57499-57969 m)	No Hits found	
SeqID 536	SA-1413.1	Contig97 (10056-12434 p)		Identities = 360/785 (45%), Positives = 496/785 (62%), Gaps = 15/785 (1%) pir  S76896 hypothetical protein - Synechocystis sp. (strain PCC 6803) dbj BAA18808.1  (D90917) hypothetical protein [Synechocystis sp.] Length = 821
SeqID 537	SA-1414.1	Contig97 (8875-9966 p)	73	Identities = 211/363 (58%), Positives = 270/363 (74%), Gaps = 9/363 (2%) splP39300 YJFR_ECOLI HYPOTHETICAL 40.1 KD PROTEIN IN AIDB-SGAT INTERGENIC REGION Length = 354
SeqID 538	SA-1415.1	Contig97 (6879-8558 p)	41	Identities = 136/511 (26%), Positives = 234/511 (45%), Gaps = 29/511 (5%) dbj BAB03939.1  (AP001507) unknown conserved protein [Bacillus halodurans]
SeqID 539	SA-1416.1	Contig97 (6052-6813 p)	68	Identities = 58/235 (24%), Positives = 104/235 (43%), Gaps = 9/235 (3%) pir] C83362 hypothetical protein PA2260 [imported] Pseudomonas aeruginosa (strain PAO1) gb AAG05648 1 AE004652_1 (AE004652) hypothetical protein [Pseudomonas aeruginosa] Length = 260
SeqID 540	SA-1417.1	Contig97 (5766-6032 p)	40	Identities = 28/89 (31%), Positives = 44/89 (48%), Gaps = 3/89 (3%) splP44991 LYXK_HAEIN PROBABLE L-XYLULOSE KINASE (L-XYLULOSE pir  H64164 hypothetical protein H11027 - Haemophilus influenzae (strain Rd KW20) gb AAC22687.1  (U32783) L-xylulose kinase (lyx) [Haemophilus influenzae Rd] Length = 485
SeqID 541	SA-1418.1	Contig97 (4513-5727 p)	48	Identities = 126/393 (32%), Positives = 212/393 (53%), Gaps = 16/393 (4%) splP37677 LYXK_ECOL! CRYPTIC L-XYLULOSE KINASE (L-XYLULOKINASE) pir  S47801 L-xylulokinase (EC 2.7.1.53) - Escherichia coli gb AAB18557.1  (U00039) No definition line found [Escherichia coli] gb AAC76604.1  (AE000435) L-xylulose kinase, cryptic [Escherichia coli K12] Length = 498
SeqID 542	SA-1419.1	Contig97 (3458-4447 p)	No Hits found	

SeqID 543	SA-1421.1	Contig97 (1985-3436 p)	40	Identities = 106/402 (26%), Positives = 196/402 (48%), Gaps = 15/402 (3%) dbj BAA75340.1  (AB011837) PTS system galactitolspecific enzyme IIC component [Bacilius halodurans] dbj BAB03909.1  (AP001507) PTS system, galactitol-specific enzyme II, C component [Bacilius halodurans]  Length = 419
SeqID 544	SA-1424.1	Contig97 (931-1887 p)	69	Identities = 123/326 (37%), Positives = 193/326 (58%), Gaps = 8/326 (2%) pir  B75057 glycerate dehydrogenase PAB2374 - Pyrococcus abyssi (strain Orsay) emb CAB50351.1  (AJ248287) GLYCERATE DEHYDROGENASE [Pyrococcus abyssi] Length = 335
SeqID 545.	SA-1425.1	Contig97 (1-912 p)	No Hits found	
SeqID 546	SA-1427.1	Contig115 (8540-9379 p)	29	Identities = 101/275 (36%), Positives = 165/275 (59%), Gaps = 22/275 (8%) gb[AAD50462.1 AF169967_5 (AF169967) BacA [Flavobacterium johnsoniae] Length = 265
SeqID 547	SA-1429.1	Contig115 (9499-10254 p)	. 80	Identities = 168/248 (67%), Positives = 205/248 (81%), Gaps = 9/248 (3%) dbjjBAA82113.1  (AB022909) negative regulator of genetic competence [Streptococcus mutans] Length = 240
SeqiD 548	SA-143.1	Contig138 (88238-89581 p)	. 40	Identities = 103/460 (22%), Positives = 191/460 (41%), Gaps = 72/460 (15%) pir  F69774 hypothetical protein ydcQ [imported] - Bacillus subtilis dbj BAA19323.1  (AB001488) SIMILAR TO ORF21 OF ENTEROCOCCUS FAECALIS TRANSPOSON TN916. [Bacillus subtilis] emb CAB12293.1  (Z99106) similar to transposon protein [Bacillus subtilis]
SeqID 549	SA-1430.1	Contig115 (10400-11416 p)	80	Identities = 267/382 (69%), Positives = 317/382 (82%)   dbj BAA82114.1  (AB022909) RgpG [Streptococcus mutans]   Length = 388

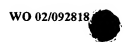


Identities = 180/250 (72%), Positives = 212/250 (84%) splP80866 v296_BACSU VEGETATIVE PROTEIN 296 (VEG296) pir] H70019 ABC transporter (ATP-binding protein) homolog yurY - Bacillus subtilis emb CAB15260.1  (299120) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis] Length = 261	Identities = 174/435 (40%), Positives = 269/435 (61%), Gaps = 15/435 (3%) dbj BAB07189.1 (AP001518) unknown conserved protein [Bacillus halodurans]  Length = 435	Identities = 241/400 (60%), Positives = 308/400 (76%), Gaps = 5/400 (1%) pir[JF70019 nifS protein homolog yurW - Bacillus subtilis emb[CAB15258.1] (299120) similar to NifS protein homolog [Bacillus subtilis] Length = 406	Identities = 72/139 (51%), Positives = 92/139 (65%) splO32163 NIFU_BACSU NIFU-LIKE PROTEIN piri E70019 nitrogen fixation protein nifU homolog yurV - Bacillus subtilis emb CAB15257.1  (299120) similar to NifU protein homolog [Bacillus subtilis]	Identities = 315/459 (68%), Positives = 385/459 (83%) pir  D70019 conserved hypothetical protein yurU - Bacillus subtilis emb CAB15256.1  (299120) similar to hypothetical proteins [Bacillus subtilis] Length = 465	Identities = 117/333 (35%), Positives = 187/333 (56%), Gaps = 8/333 (2%) emb CAA60585.1  (X87105) penicillin binding protein 4 [Staphylococcus aureus] emb CAA60582.1  (X87104) penicillin binding protein 4 [Staphylococcus aureus]	Identities = 203/415 (48%), Positives = 300/415 (71%), Gaps = 6/415 (1%) emb CAA67776.1  (X99400) D,D-carboxypeptidase [Streptococcus pneumoniae] Length = 413
77	62	74	58	79	45	71
Contig115 (11581-12351 p)	Contig115 (12388-13650 p)	Contig115 (13652-14884 p)	Contig115 (14871-15314 p)	Contig115 (15414-16832 p)	Contig115 (16904-18031 m)	Contig115 (18244-19479 m)
SA-1431.1	SA-1432.1	SA-1433.1	SA-1434.1	SA-1435.1	SA-1436.1	SA-1437.2
SeqID 550	SeqID 551	SeqID 552	SeqID 553	SeqID 554	SeqID 555	SeqID 556

SeqID 557	SA-1438.2	Contig139 (86184-86444 m)	17	Identities = 61/81 (75%), Positives = 71/81 (87%), Gaps = 1/81 (1%) gb AAF80389.1 AF160251_3 (AF160251) ribosomal protein L31 [Listeria innocua]
SeqID 558	SA-1439.1	Contig139 (84551-86071 p)	61	Identities = 257/429 (59%), Positives = 312/429 (71%), Gaps = 7/429 (1%) pir  T46756 Zn-binding lipoprotein adcA [imported] - Streptococcus pneumoniae (fragment) emb CAA96185.1  (Z71552) AdcA protein [Streptococcus pneumoniae] Length = 423
SeqID 559	SA-144.1	Contig138 (87835-88260 p)	No Hits found	
SeqID 560	SA-1440.1	Contig139 (83012-84409 p)	65	Identities = 218/473 (46%), Positives = 310/473 (65%), Gaps = 14/473 (2%) emb CAA86210.1  (Z38063) dipeptidase [Lactobacillus helveticus]
SeqID 561	SA-1441.1	Contig139 (82086-82997 p)	82	Identities = 209/303 (68%), Positives = 260/303 (84%) emb CAB96619.1  (AJ400630) hypothetical protein [Streptococcus pneumoniae bacteriophage MM1] emb CAB96622.1  (AJ400631) hypothetical protein [Streptococcus pneumoniae] Length = 303
SeqID 562	SA-1442.2	Contig139 (81115-82089 p)	98	Identities = 254/321 (79%), Positives = 286/321 (88%), Gaps = 1/321 (0%) emb[CAB96620.1  (AJ400630) hypothetical protein [Streptococcus pneumoniae bacteriophage MM1] emb[CAB96623.1  (AJ400631) hypothetical protein [Streptococcus pneumoniae] Length = 325
SeqID 563	SA-1445.2	Contig139 (80228-81118 p)	89	Identities = 165/291 (56%), Positives = 208/291 (70%), Gaps = 2/291 (0%) sp 006973 YVCJ_BACSU HYPOTHETICAL 33.9 KD PROTEIN IN CRH-TRXB INTERGENIC REGION pir  H70031 conserved hypothetical protein yvcJ - Bacillus subtilis emb CAB08057.1  (294043) hypothetical protein [Bacillus subtilis] emb CAB15482.1  (299121) similar to hypothetical proteins [Bacillus subtilis]

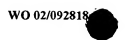


SeqID 564	SA-1446.1	Contig139 (79661-80074 m)	98	Identities = 36/134 (26%), Positives = 57/134 (41%), Gaps = 7/134 (5%) emb CAB62728.1  (AL133423) hypothetical protein SC4A7.24c [Streptomyces coelicolor A3(2)] Length = 139
SeqID 565	SA-1447.1	Contig139 (78432-79415 m)	46	Identities = 97/305 (31%), Positives = 160/305 (51%), Gaps = 10/305 (3%) splP33022 YEIK_ECOLI HYPOTHETICAL 33.7 KD PROTEIN IN NFO-FRUA INTERGENIC REGION pir  A64985 hypothetical 33.7 kD protein in nfo-fruA intergenic region - Escherichia coli (strain K-12) gb AAA60514.1 (U00007) yeiK [Escherichia coli] gb AAC75223.1 (AE000305) orf, hypothetical protein [Escherichia coli K12] prf  2014253BH yeiK gene [Escherichia coli]
SeqID 566	SA-1448.1	Contig139 (77878-78435 m)	38	Identities = 39/135 (28%), Positives = 72/135 (52%), Gaps = 4/135 (2%) ref[NP_053012.1] hypothetical protein [Plasmid pNZ4000] gb[AAD40355.1] (AF035485) hypothetical protein [Plasmid pNZ4000]
SeqID 567	SA-1449.1	Contig139 (76492-77838 p)	69	Identities = 252/442 (57%), Positives = 316/442 (71%), Gaps = 15/442 (3%) dbj BAB05415.1  (AP001512) asparaginyl-tRNA synthetase [Bacillus halodurans] Length = 430
SeqID 568	SA-145.1	Contig138 (87488-87769 p)	No Hits found	
SeqID 569	SA-1450.1	Contig139 (75278-76471 p)	75	Identities = 270/391 (69%), Positives = 314/391 (80%) gb AAF12702.1 AF035157_1 (AF035157) aspartate aminotransferase [Lactococcus lactis] Length = 393
SeqID 570	SA-1451.1	Contig139 (72727-75192 p)	47	Identities = 251/927 (27%), Positives = 398/927 (42%), Gaps = 145/927 (15%) dbjjBAB05410.1 (AP001512) ATP-dependent DNA helicase [Bacillus halodurans]  Length = 942
SeqID 571	SA-1453.2	Contig139 (71813-72577 p)	69	Identities = 162/254 (63%), Positives = 188/254 (73%), Gaps = 2/254 (0%) gb AAC48769.1  (U71200) acetoin reductase [Bos taurus]



<u></u>	Ι		1	1		1		
Identities = 53/112 (47%), Positives = 78/112 (69%), Gaps = 1/112 (0%) gb AAC23746.1  (AF052209) competence protein [Streptococcus pneumoniae]	Identities = 142/468 (30%), Positives = 220/468 (46%), Gaps = 62/468 (13%) pir[ F82995 glutamatecysteine ligase PA5203 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG08588.1 AE004933_4 (AE004933) glutamatecysteine ligase [Pseudomonas aeruginosa] Length = 527	Identities = 107/223 (47%), Positives = 148/223 (65%), Gaps = 7/223 (3%) pir  H81283 helix-turn-helix containing protein Cj1387c [imported] - Campylobacter jejuni (strain NCTC 11168) emb CAB73814.1  (AL139078) helix-turn-helix containing protein [Campylobacter jejuni] Length = 218		Identities = 225/341 (65%), Positives = 279/341 (80%) pir  A43577 regulatory protein pfoR - Clostridium perfringens Length = 343	Identities = 320/427 (74%), Positives = 378/427 (87%) pir  A42280 adenylosuccinate synthase (EC 6.3.4.4) purA - Bacillus subtilis dbj BAA05174.1  (D26185) adenylosuccinate synthetase [Bacillus subtilis] emb CAB16079.1  (299124) adenylosuccinate synthetase [Bacillus subtilis] Length = 430			Identities = 184/465 (39%), Positives = 279/465 (59%), Gaps = 25/465 (5%) splP39301 SGAT_ECOLI PUTATIVE TRANSPORT PROTEIN SGAT pir  D65230 hypothetical 52.9 kD protein in aidB rpsF intergenic region - Escherichia coli (strain K-12) gb AAC77150.1  (AE000491) orf, hypothetical protein [Escherichia coli K12] Length = 484
09	29	64	No Hits found	77	87	No Hits found	No Hits found	55
Contig95 (12824-13120 m)	Contig95 (10627-12858 p)	Contig95 (9882-10556 p)	Contig138 (86863-87177 m)	Contig95 (8498-9520 m)	Contig95 (6835-8127 m)	Contig95 (6339-6812 p)	Contig95 (5563-6180 m)	Contig95 (3956-5395 m)
SA-1456.1	SA-1457.1	SA-1458.1	SA-146.1	SA-1460.1	SA-1462.1	SA-1463.1	SA-1465.1	SA-1466.1
SeqID 572	SeqID 573	SeqID 574	SeqID 575	SeqID 576	SeqID 577	SeqID 578	SeqID 579	SeqID 580

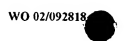




SeqID 586	SA-1471.1	Contig95 (735-1451 m)	73	Identities = 143/229 (62%), Positives = 176/229 (76%), Gaps = 2/229 (0%) gb AAD45716.1 AF160811_4 (AF160811) L-ribulose 5-phosphate 4-epimerase [Bacillus stearothermophilus] Length = 228
SeqID 587	SA-1472.1	Contig95 (77-724 m)	09	Identities = 124/214 (57%), Positives = 157/214 (72%) splQ58370 TAL_METJA TRANSALDOLASE-LIKE PROTEIN pir  H64419 transaldolase (EC 2.2.1.2) - Methanococcus jannaschii gb AAB98962.1  (U67539) transaldolase [Methanococcus jannaschii] Length = 217
SeqID 588	SA-1473.2	1	No Hits found	
SeqID 589 SeqID 590	SA-1474.1 SA-1475.1	Contig94 (11451-11573 m) Contig94 (10797-11117 m)	No Hits found	
SeqID 591	SA-1477.1	1	48	Identities = 59/185 (31%), Positives = 96/185 (51%), Gaps = 12/185 (6%) pir  A82466 hypothetical protein VCA0380 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF96286.1  (AE004374) hypothetical protein [Vibrio cholerae] Length = 201
SeqID 592	SA-1478.1	Contig94 (8817-9236 m)	62	Identities = 49/147 (33%), Positives = 98/147 (66%), Gaps = 2/147 (1%) pir  A69849 hypothetical protein yjdF - Bacilius subtilis emb CAB13060.1  (299110) yjdF [Bacilius subtilis] Length = 160
SeqID 593	SA-1479.1	Contig94 (8138-8617 m)	58	Identities = 61/152 (40%), Positives = 95/152 (62%) gb AAD50427.1 AF161700_2 (AF161700) ComX1 [Streptococcus pneumoniae] gb AAD50429.1 AF161701_2 (AF161701) ComX2 [Streptococcus pneumoniae] Length = 159
SeqID 594	SA-148.1	Contig138 (85248-85790 m)	46	Identities = 55/175 (31%), Positives = 93/175 (52%), Gaps = 6/175 (3%) splP05332 YP20_BACLI HYPOTHETICAL P20 PROTEIN pir  S00875 hypothetical protein P20 - Bacillus licheniformis emb CAA30415.1  (X07542) P20 (AA 1-178) [Bacillus licheniformis] Length = 178



SeqID 595	SA-1480.1	Contig94 (7324-8016 m)	14	Identities = 57/226 (25%), Positives = 97/226 (42%), Gaps = 22/226 (9%) ref[NP_014926.1] Yor283wp [Saccharomyces cerevisiae] pir[ S67185 hypothetical protein YOR283w - yeast (Saccharomyces cerevisiae) emb CAA61787.1  (X89633) hypothetical phosphoglycerate mutase [Saccharomyces cerevisiae] emb CAA99510.1  (Z75191) ORF YOR283w [Saccharomyces cerevisiae] Length = 230
Seq1D 596	SA-1481.1	Contig94 (6575-7249 m)	50	Identities = 108/169 (63%), Positives = 139/169 (81%) gb AAD00280.1  (U78599) putative D,D-carboxypeptidase [Streptococcus mutans] Length = 173
SeqID 597	SA-1482.1	Contig94 (6003-6425 m)	84	Identities = 65/153 (42%), Positives = 94/153 (60%), Gaps = 4/153 (2%) gb AAD00279.1 (U78599) putative N-acetylmuramidase [Streptococcus mutans]
SeqID 598	SA-1483.1	Contig94 (4826-5860 m)	80	Identities = 233/344 (67%), Positives = 294/344 (84%) splO06940JHRCA_STRMU HEAT-INDUCIBLE TRANSCRIPTION REPRESSOR HRCA Length = 344
SeqID 599	SA-1484.1	Contig94 (4251-4784 m)	. 78	Identities = 129/188 (68%), Positives = 153/188 (80%), Gaps = 2/188 (1%) pir  S39341 grpE protein - Lactococcus lactis Length = 190
SeqID 600	SA-1486.1	Contig94 (2241-4070 m)	. 26	Identities = 609/609 (100%), Positives = 609/609 (100%) splP95693 DNAK_STRAG DNAK PROTEIN (HEAT SHOCK PROTEIN 70) (HSP70) gb AAB39219.1  (U72719) heat shock protein 70 (Streptococcus agalactiae) Length = 609
SeqID 601	SA-1487.1	Contig94 (813-1952 m)	693	Identities = 328/377 (87%), Positives = 356/377 (94%), Gaps = 1/377 (0%) dbj BAB16032.1  (AB030809) Streptococcus pneumoniae DnaJ protein homologue [Streptococcus pyogenes] Length = 378
SeqID 602	SA-1488.1	Contig94 (495-827 p)	No Hits found	
SeqID 603	SA-1489.1	Contig94 (1-522 p)	62	Identities = 122/164 (74%), Positives = 138/164 (83%) splQ07211[SCRK_STRMU FRUCTOKINASE db] BAA02467.1  (D13175) fructokinase [Streptococcus mutans] Length = 293



SeqID 604	SA-149.1	Contig138 (84525-85205 m)	2	Identities = 110/230 (47%). Positives = 154/230 (66%). Gaps = 3/230 (1%) ref[NP_012308 1] Yir042cp [Saccharomyces cerevisiae] spiP40585/YW2_YEAST HYPOTHETICAL 27.4 KD PROTEIN IN HYR1 3 REGION pir[]S50347 hypothetical protein YIR042c · yeast (Saccharomyces cerevisiae) emb[CAA87001.1] (246902) unknown [Saccharomyces cerevisiae] Length = 236
SeqID 605	SA-1490.2	Contig134 (49677-50102 p)	83	Identities = 118/139 (84%), Positives = 129/139 (91%) splP36254[RL11_STACA 50S RIBOSOMAL PROTEIN L11 pir  S38871 ribosomal protein L11 - Staphylococcus carnosus emb CAA53739.1  (X76134) L11 protein [Staphylococcus carnosus] Length = 140
SeqID 606	SA-1491.1	Contig134 (48084-49466 p)	43	Identities = 118/405 (29%), Positives = 199/405 (49%), Gaps = 9/405 (2%) pir[ F70705 hypothetical protein Rv2333c - Mycobacterlum tuberculosis (strain H37RV) emb CAB02058.1  (279702) hypothetical protein Rv2333c [Mycobacterium tuberculosis] Length = 537
SeqID 607	SA-1492.1	Contig134 (46862-48076 p)	99	Identities = 201/395 (50%), Positives = 274/395 (68%), Gaps = 5/395 (1%) gb AAF36227.1 AF168363_2 (AF168363) aminoacylase [Lactococcus lactis] Length = 398
SeqID 608	SA-1493.1	Contig134 (45698-46582 m)	40	Identities = 72/259 (27%), Positives = 127/259 (48%), Gaps = 7/259 (2%) refINP_043632.1 former trsE (rbcR homolog) [Odontella sinensis] splP49518 YC30_ODOSI HYPOTHETICAL TRANSCRIPTIONAL REGULATOR YCF30 pir  S78291 probable transcription regulator ycf30 - Odontella sinensis chloroplast emb CAA91664.1 (267753) former trsE (rbcR homolog) [Odontella sinensis]
SeqID 609	SA-1494.1	Contig134 (45284-45646 m)	No Hits found	



SeqID 610	SA-1495.1	Contig134 (43540-45255 p)	56	Identities = 203/580 (35%), Positives = 325/580 (56%), Gaps = 50/580 (8%) pir  E64556 para-aminobenzoate synthetase - Helicobacter pylori (strain 26695) gb AAD07357.1  (AE000547) para-aminobenzoate synthetase (pabB) [Helicobacter pylori 26695] Length = 559
SeqID 611	SA-1498.1	Contig134 (41017-43458 p)	62	Identities = 353/828 (42%), Positives = 511/828 (61%), Gaps = 70/828 (8%) sp P21458 SP3E_BACSU STAGE III SPORULATION PROTEIN E pir  S09411 DNA translocase spollIE - Bacilius subtilis emb CAB13553.1  (Z99112) DNA translocase [Bacilius subtilis]
SeqID 612	SA-1499.1	Contig134 (40039-40842 m)	44	Identities = 88/224 (39%), Positives = 123/224 (54%), Gaps = 46/224 (20%) pir  T41399 probable cyclophilin-related peptidyl prolyl cis-trans isomerase - fission yeast (Schizosaccharomyces pombe) emb CAA19257.1  (AL023704) putative Cyclophilin-type peptidyl-prolyl cis-trans isomerase protein [Schizosaccharomyces pombe] Length = 610
SeqID 613	SA-15.1	Contig137 (29069-31870 p)	22	Identities = 152/405 (37%), Positives = 214/405 (52%), Gaps = 26/405 (6%) piri[A43607 cell surface antigen SpaA precursor - Streptococcus sobrinus (strain MT3791) dbj[BAA14368.1] (D90354) surface protein antigen precursor [Streptococcus sobrinus] Length = 1566
SeqID 614	SA-150.1	Contig138 (83318-84265 p)	No Hits found	
SeqID 615	SA-1500.1	Contig134 (39154-39987 p)	06	Identities = 224/275 (81%), Positives = 255/275 (92%) gb AAD56938.1/AF180520_3 (AF180520) integral membrane protein MtsC [Streptococcus pyogenes] Length = 284
SeqID 616	SA-1501.1	Contig134 (38436-39152 p)	72	Identities = 143/238 (60%), Positives = 186/238 (78%), Gaps = 2/238 (0%) gb AAD56937.1 AF180520_2 (AF180520) ATP-binding protein MtsB [Streptococcus pyogenes] Length = 241

SeqID 617	SA-1502.2	Contig134 (37339-38265 p)	- 98	Identities = 240/308 (77%), Positives = 277/308 (89%), Gaps = 1/308 (0%) gbjAAD56936.1 AF180520_1 (AF180520) lipoprotein MtsA [Streptococcus pyogenes] Length = 310
SeqID 618	SA-1503.2	Contig108 (5713-10425 m)	99	Identities = 795/1596 (49%), Positives = 1056/1596 (65%), Gaps = 39/1596 (2%) gb AAG09771.1 AF243528_1 (AF243528) cell envelope proteinase [Streptococcus thermophilus] Length = 1585
SeqID 619	SA-1504.1	Contig108 (4794-5480 m)	20	Identities = 87/220 (39%), Positives = 124/220 (55%), Gaps = 4/220 (1%) dbj BAB05663.1  (AP001513) two-component response regulator [Bacillus halodurans]
SeqID 620	SA-1506.1	Contig108 (3409-4665 m)	40	Identities = 108/379 (28%), Positives = 193/379 (50%), Gaps = 33/379 (8%) pir  D70045 two-component sensor histidine kinase homolog yvqB - Bacillus subtilis emb CAB15292.1  (Z99120) similar to two-component sensor histidine kinase [YvqA] [Bacillus subtilis] emb CAA11751.1  (AJ223978) putative sensory protein kinase, YvqB [Bacillus subtilis]
SeqID 621	SA-1507.1	Contig108 (2708-3334 m)	33	Identities = 38/144 (26%), Positives = 69/144 (47%), Gaps = 10/144 (6%) ref[XP_005848.1  KIAA1074 protein [Homo sapiens] Length = 1709
SeqID 622	SA-151.1	Contig138 (82830-83321 p)	42	Identities = 40/141 (28%), Positives = 75/141 (52%), Gaps = 3/141 (2%) dbj BAB05334.1  (AP001512) RNA polymerase sigma factor Y [Bacillus halodurans] Length = 176
SeqID 623	SA-1511.1	Contig108 (120-2621 p)	78	Identities = 569/835 (68%), Positives = 666/835 (79%), Gaps = 42/835 (5%) splP36430 SYL_BACSU LEUCYL-TRNA SYNTHETASE (LEUCINETRNA LIGASE) (LEURS) pirj D69650 leucinetRNA ligase (EC 6.1.1.4) - Bacillus subtilis gb AAC00259.1  (AF008220) leucine tRNA synthetase [Bacillus subtilis] emb CAB15010.1  (299119) leucyl-tRNA synthetase [Bacillus subtilis]

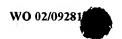


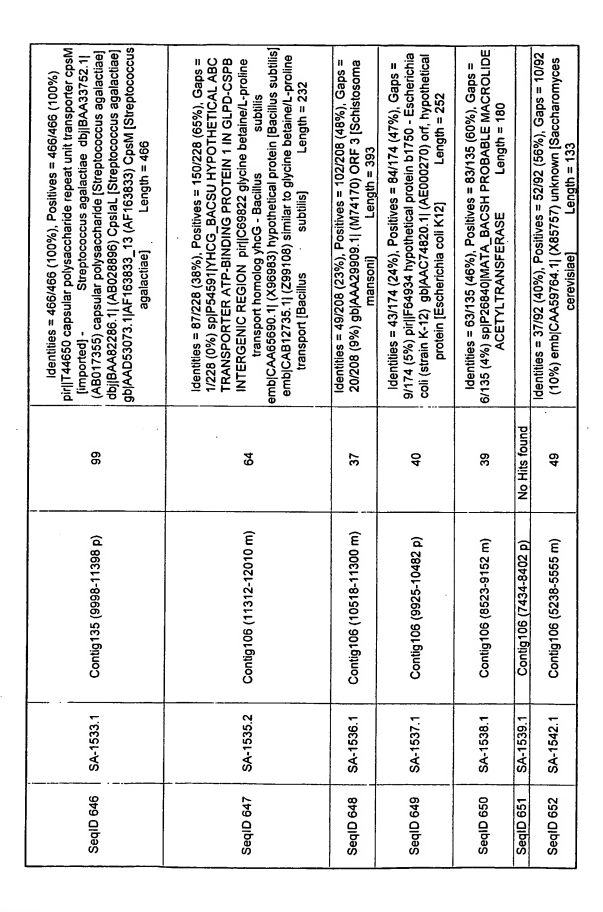
SeqID 624	SA-1512.2	Contig138 (776-1330 m)	17	Identities = 107/185 (57%), Positives = 140/185 (74%), Gaps = 3/185 (1%) emb CAB91549.1  (AJ249134) hypothetical protein [Lactococcus lactis] Length = 185
SeqID 625	SA-1513.1	Contig138 (1407-2072 m)	51	Identities = 76/230 (33%), Positives = 118/230 (51%), Gaps = 11/230 (4%) splP39147 CMF3_BACSU COMF OPERON PROTEIN 3 pir[ 140389 hypothetical protein F3 - Bacillus subtilis pir  S77621 late competence gene comFC - Bacillus subtilis emb CAA79228.1  (218629) F3 [Bacillus subtilis] gb AAC44942.1  (U56901) involved in transformation [Bacillus subtilis] emb CAB15562.1  (299122) late competence gene [Bacillus subtilis]
SeqID 626	SA-1514.1	Contig138 (2838-3017 p)	No Hits found	
SeqID 627	SA-1515.1	Contig138 (2072-3361 m)	92	Identities = 160/405 (39%), Positives = 228/405 (55%), Gaps = 20/405 (4%) splP39145 CMF1_BACSU COMF OPERON PROTEIN 1 pir  40387 hypothetical protein F1 - Bacillus subtilis pir  G69602 late competence protein required for DNA uptake comFA - Bacillus subtilis emb CAA79226.1  (Z18629) F1 [Bacillus subtilis] gb AAC44940.1  (U56901) involved in transformation [Bacillus subtilis] emb CAB15564.1  (Z99122) late competence protein [Bacillus subtilis]
SeqID 628	SA-1516.1	Contig138 (3417-4061 p)	58	Identities = 96/204 (47%), Positives = 127/204 (62%) dbj BAB07349.1  (AP001519) unknown conserved protein [Bacillus halodurans] Length = 213
SeqID 629	SA-1517.1	Contig138 (4152-5078 p)	88	Identities = 239/306 (78%), Positives = 274/306 (89%) dbj BAA88310.1 .(AB028865) O-acetylserine lyase [Streptococcus suis] Length = 308
SeqID 630	SA-1518.1	Contig138 (5230-6324 m)	73	Identities = 199/363 (54%), Positives = 272/363 (74%), Gaps = 2/363 (0%) splP32816 GLDA_BACST GLYCEROL DEHYDROGENASE (GLDH) pir JQ1474 glycerol dehydrogenase (EC 1.1.1.6) [validated] - Bacillus stearothermophilus gblAAA22477.1  (M65289) glycerol dehydrogenase [Bacillus stearothermophilus] Length = 370





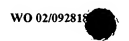
SeqID 638	SA-1525.1	Contig135 (16949-17587 m)	29	Identities = 134/213 (62%), Positives = 169/213 (78%) splQ54916 YPAE_STRPN HYPOTHETICAL 23.0 KD PROTEIN IN PARE 5 REGION (ORF2) emb CAA91549.1  (Z67739) unidentified [Streptococcus pneumoniae] Length = 213
SeqID 639	SA-1526.1	Contig135 (16230-16883 p)		Identities = 217/217 (100%), Positives = 217/217 (100%) splQ9XDS8IUNG_STRAG URACIL-DNA GLYCOSYLASE (UDG) dbj BAA82292.1  (AB028896) Uracil DNA glycosylase [Streptococcus agalactiae] Length = 217
SeqID 640	SA-1527.1	Contig135 (15646-16131 p)	96	Identities = 161/161 (100%), Positives = 161/161 (100%) dbj BAA82291.1  (AB028896) ORF1 [Streptococcus agalactiae] Length = 161
SeqiD 641	SA-1528.1	Contig135 (14291-15532 p)	86	Identities = 413/413 (100%), Positives = 413/413 (100%) dbj BAA82290.1  (AB028896) NeuA [Streptococcus agalactiae] Length = 413
SeqID 642	SA-153.1	Contig138 (80519-81616 p)	30	Identities = 64/187 (34%), Positives = 112/187 (59%), Gaps = 1/187 (0%) pir  T36287 probable integral membrane protein - Streptomyces coelicolor emb CAB42664.1  (AL049819) putative integral membrane protein [Streptomyces coelicolor A3(2)]
SeqID 643	SA-1530.1	Contig135 (13651-14280 p)	86	Identities = 209/209 (100%), Positives = 209/209 (100%) dbj BAA82289.1  (AB028896) NeuD [Streptococcus agalactiae] Length = 209
SeqID 644	SA-1531.1	Contig135 (12500-13654 p)	96	Identities = 384/384 (100%), Positives = 384/384 (100%) dbj BAA82288.1  (AB028896) NeuC [Streptococcus agalactiae] Length = 384
SeqID 645	SA-1532.1	Contig135 (11398-12423 p)	26	Identities = 341/341 (100%), Positives = 341/341 (100%) dbj BAA33753.1  (AB017355) neuB [Streptococcus agalactiae] dbj BAA82287.1  (AB028896) NeuB [Streptococcus agalactiae] Length = 341







Identities = 410/922 (44%), Positives = 609/922 (65%), Gaps = 22/922 (2%) pir  C69069 cation-transporting P-ATPase PacL - Methanobacterium thermoautotrophicum (strain Delta H) gb AAB85991.1  (AE000912) cation-transporting P-ATPase PacL [Methanobacterium thermoautotrophicum] Length = 910	Identities = 85/132 (64%), Positives = 105/132 (79%) emb CAB46979.1  (AJ243482) CSRA protein [Enterococcus faecalis] Length = 168	Identities = 48/152 (31%), Positives = 75/152 (48%), Gaps = 1/152 (0%) dbj BAB05127.1  (AP001511) unknown [Bacilius halodurans] Length = 156	Identities = 108/211 (51%), Positives = 149/211 (70%) dbj BAB06554.1  (AP001516) unknown conserved protein [Bacillus halodurans] Length = 215	Identities = 76/279 (27%), Positives = 133/279 (47%), Gaps = 4/279 (1%) splP25146 INLA_LISMO INTERNALIN A PRECURSOR pir  S37387 internalin A precursor - Listeria monocytogenes Length = 800	Identities = 86/232 (37%), Positives = 137/232 (58%), Gaps = 3/232 (1%) gb AAF50837.1  (AE003568) CG1718 gene product [Drosophila melanogaster] Length = 1713	Identities = 41/110 (37%), Positives = 56/110 (50%), Gaps = 4/110 (3%) gb AAD33086.1 AF071083_1 (AF071083) fibronectinbinding protein I [Streptococcus pyogenes] Length = 1161		Identities = 58/135 (42%), Positives = 86/135 (62%), Gaps = 2/135 (1%) gb AAC13546.1  (AF019629) putative fimbria-
1de 22 22 Me 65 9b/A	67 e	1 10	99	4L	1d 1d 3/2	lo 14/11 bing	No Hits found	10 10 10 10 10 10 10 10 10 10 10 10 10 1
							No Hit	
Contig106 (4507-7299 p)	Contig106 (3921-4355 p)	Contig106 (3451-3921 p)	Contig106 (2795-3448 p)	Contig106 (26-2689 p)	Contig138 (79800-80516 p)	Contig139 (128987-131698 p)	Contig139 (131737-131979 p)	Contig139 (132099-132542 p)
SA-1543.1	SA-1544.1	SA-1545.1	SA-1546.1	SA-1547.1	SA-155.1	SA-1551.2	SA-1552.1	SA-1554.1
SeqID 653	SeqID 654	SeqID 655	SeqID 656	SeqID 657	SeqID 658	SeqID 659	SeqID 660	SeqID 661



SeqID 662	SA-1555.1	Contig139 (132758-133396 p)	19	Identities = 30/96 (31%), Positives = 46/96 (47%), Gaps = 9/96 (9%) gb AAD33086.1 AF071083_1 (AF071083) fibronectinbinding protein I [Streptococcus pyogenes] Length = 1161
SeqID 663	SA-1556.1	Contig 139 (133726-133872 p)	No Hits found	
SeqID 664	SA-1558.1	Contig139 (134494-134670 p)	46	Identities = 23/67 (34%), Positives = 37/67 (54%), Gaps = 2/67 (2%) gb AAB00100.1  (L36660) unknown [Streptococcus pneumoniae] emb CAA84075.1  (Z34303) hypothetical protein [Streptococcus pneumoniae] Length = 194
SeqID 665	SA-156.1	Contig138 (78853-79536 p)	38	Identities = 45/205 (21%), Positives = 93/205 (44%), Gaps = 10/205 (4%) dbj BAB04126.1  (AP001508) unknown conserved protein in others [Bacillus halodurans] Length = 228
SeqID 666	SA-1560.1	Contig139 (135988-136284 p)	No Hits found	
SeqID 667	SA-1562.1	Contig139 (136414-137496 p)	31	Identities = 72/279 (25%), Positives = 125/279 (43%), Gaps = 25/279 (8%) dbj BAA24012.1  (AB009635) Fmt [Staphylococcus aureus] Length = 397
SeqID 668	SA-1563.1	Contig139 (137702-138688 p)	58	Identitles = 141/331 (42%), Positives = 201/331 (60%), Gaps = 34/331 (10%) pir  G75548 ABC transporter, ATP-binding protein - Deinococcus radiodurans (strain R1) gb AAF09790.1 AE001882_8 (AE001882) ABC transporter, ATP-binding protein {Deinococcus radiodurans} Length = 354
SeqID 669	SA-1564.1	Contig139 (138691-139509 p)	42	Identities = 60/271 (22%), Positives = 118/271 (43%), Gaps = 13/271 (4%) emb CAB88837.1  (AL353832) putative integral membrane protein. [Streptomyces coelicolor A3(2)]  Length = 268



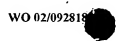
	1-140296 p) 44	Contig139 (139511-140296 p) 44	
Identities = 25/84 (29%), Positives = 45/84 (52%), Gaps = 6/84 (7%) gbjAAD31042 1jAF144646_1 (AF144646) heat shock protein 70 [Crassostrea gigas]	41		12896-13207 m) 41
	3-14211 p) 67	Contig128 (13333-14211 p) 67	13333-14211 p)
Identities = 67/133 (50%), Positives = 86/133 (64%), Gaps = 16/133 (12%) dbj BAB04125.1  (AP001508) unknown conserved protein in others [Bacillus halodurans] Length = 174	72		78521-78832 p) 72
Identities = 120/203 (59%), Positives = 145/203 (71%), Gaps = 9/203 (4%) sp O07344 LEP_STRPN SIGNAL PEPTIDASE I (SPASE I) (LEADER PEPTIDASE I) gb AAB69116.1  (U90721) signal peptidase I [Streptococcus pneumoniae] Length = 204	71		14227-14820 p) 71
Identities = 454/835 (54%); Positives = 600/835 (71%), Gaps = 37/835 (4%) gb AAK05838.1 AE006404_5 (AE006404) exodeoxyribonuclease V alpha chain (EC 3.1.11.5) [Lactococcus lactis subsp. lactis] Length = 834	, 73		
·	3-17965 p) 42	Contig128 (17483-17965 p) 42	17483-17965 p)



SeqID 677	SA-1574.1	Contig128 (18036-19055 m)		Identities = 136/349 (38%), Positives = 210/349 (59%), Gaps = 14/349 (4%) pir  G82093 DNA-damage-inducible protein P VC2287 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF95431.1  (AE004300) DNA-damage-inducible protein P [Vibrio cholerae] Length = 360
SeqID 678	SA-1577.1	Contig128 (19315-21627 p)	95	Identities = 708/770 (91%), Positives = 750/770 (96%) splQ59934 PFL_STRMU FORMATE ACETYLTRANSFERASE (PYRUVATE FORMATE-LYASE) dbj BAA09085.1  (D50491) Pyruvate formate-lyase [Streptococcus mutans] Length = 775
SeqID 679	SA-1578.1	Contig128 (21728-22102 p)	SS	Identities = 53/124 (42%), Positives = 76/124 (60%), Gaps = 2/124 (1%) splQ46604 FMNB_DESVM FMN-BINDING PROTEIN pdb 1FLM A Chain A, Dimer Of Fmn-Binding Protein From Desulfovibrio Vulgaris (Miyazaki F) pdb 1FLM B Chain B, Dimer Of Fmn-Binding Protein From Desulfovibrio Vulgaris (Miyazaki F) pdb 1AXJ  Fmn-Binding Protein From Desulfovibrio Vulgaris (Miyazaki F), Nmr, 20 Structures db BAA25177.1  (D21804) FMN-binding protein [Desulfovibrio vulgaris]
SeqID 680	SA-1579.1	Contig128 (22137-23081 m)	44	Identities = 88/323 (27%), Positives = 142/323 (43%), Gaps = 51/323 (15%) emb CAC16441.1  (AL450165) putative esterase [Streptomyces coelicolor] Length = 353
SeqID 681	SA-1580.1	Contig128 (23063-23818 m)	55	Identities = 92/240 (38%), Positives = 143/240 (59%), Gaps = 11/240 (4%) gb AAD37110.1  (AF112358) C3-degrading proteinase [Streptococcus pneumoniae] Length = 241
SeqID 682	SA-1582.1	Contig128 (23943-24839 p)	21	Identities = 43/180 (23%), Positives = 71/180 (38%), Gaps = 16/180 (8%) dbj BAB07173.1  (AP001518) unknown conserved protein [Bacillus halodurans]
SeqID 683	SA-1584.3	Contig128 (24918-25745 m)	77	Identities = 186/290 (64%), Positives = 228/290 (78%), Gaps = 10/290 (3%) gb AAK06146.1 AE006435_4 (AE006435)   transporter [Lactococcus lactis subsp. lactis]   Length = 289

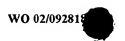


SeqID 684	SA-1585.3	Contig128 (25935-26387 m)	09	Identities = 58/144 (40%), Positives = 92/144 (63%)  gb[AAK05931.1;AE006413_7 (AE006413) conserved hypothetical protein [Lactococcus lactis subsp. lactis] Length = 145
SeqID 685	SA-1586.2	Contig128 (26405-27607 m)	48	Identities = 106/401 (26%), Positives = 199/401 (49%), Gaps = 21/401 (5%) pir B70065 antibiotic resistance protein homolog ywoG - Bacillus subtilis emb CAB05383.1  (Z82987) unknown similar to quinolon resistance protein NorA [Bacillus subtilis] emb CAB15662.1  (Z99122) similar to antibiotic resistance protein [Bacillus subtilis]
SeqID 686	SA-1587.2	Contig139 (97176-98552 p)	78	Identities = 280/458 (61%), Positives = 363/458 (79%), Gaps = 3/458 (0%) gb AAD47594.1 AF140784_3 (AF140784) Vexp3 [Streptococcus pneumoniae] Length = 459
SeqID 687	SA-1588.1	Contig139 (96520-97176 p)	76	Identities = 142/207 (68%), Positives = 169/207 (81%) gb AAD47593.1 AF140784_2 (AF140784) Vexp2 [Streptococcus pneumoniae] Length = 215
SeqID 688	SA-1589.1	Contig139 (95233-96510 p)	63	Identities = 165/425 (38%), Positives = 271/425 (62%), Gaps = 4/425 (0%) gblAAD47592.1IAF140784_1 (AF140784) Vexp1 [Streptococcus pneumoniae] Length = 425
SeqID 689	SA-159.1	Contig138 (77622-78329 m)	32	Identities = 44/150 (29%), Positives = 80/150 (53%), Gaps = 14/150 (9%) splP16396 SUBE_BACSU MINOR EXTRACELLULAR PROTEASE EPR PRECURSOR pir  SUBSMP serine proteinase (EC 3.4.21) epr precursor - Bacillus subtilis emb CAA37392.1  (X53307) prepropeptide (AA-27 to 618) [Bacillus subtilis] gb AAA22423.1  (M22407) extracellular protease precursor [Bacillus subtilis] emb CAA51571.1  (X73124) ipa-15r epr [Bacillus subtilis] emb CAB15866.1  (Z99123) extracellular serine protease [Bacillus subtilis]
SeqID 690	SA-1590.1	Contig139 (94330-94491 m)	No Hits found	





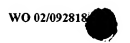
SeqID 696	SA-1597.1	Contig139 (91399-91746 p)	78	Identities = 110/115 (95%), Positives = 112/115 (96%) spl034031 RL19_STRTR 50S RIBOSOMAL PROTEIN L19 gb AAC01534.1  (U88973) ribosomal protein L19 [Streptococcus thermophilus] Length = 115
SeqID 697	SA-1598.1	Contig139 (89623-90819 p)		Identities = 131/393 (33%), Positives = 206/393 (52%), Gaps = 20/393 (5%) pir  T44296 hypothetical protein [imported] - Bacillus halodurans dbj BAA75315.1  (AB011836) similar to Bordetella paraperlussis transposase for insertion sequence element(27 -identity) [Bacillus halodurans] dbj BAB04382.1  (AP001509) unknown conserved protein in others [Bacillus halodurans] Length = 424
SeqID 698	SA-1599.1	Contig139 (89355-89630 p)	52	Identities = 24/89 (26%), Positives = 50/89 (55%), Gaps = 2/89 (2%) pir  G71950 hypothetical protein jhp0276 - Helicobacter pylori (strain J99) gb AAD05867.1  (AE001465) putative [Helicobacter pylori J99] Length = 100
SeqID 699	SA-16.1	Contig137 (28276-28605 m)	41	Identities = 35/103 (33%), Positives = 46/103 (43%), Gaps = 14/103 (13%) gb[AAC02237.1 (U72957) merozoite surface protein 2 [Plasmodium falciparum] Length = 176
SeqID 700	SA-160.2	Contig138 (76196-77368 m)	71	Identities = 236/387 (60%), Positives = 281/387 (71%), Gaps = 10/387 (2%) gb AAK05433.1 AE006366_2 (AE006366) N-acetylglucosamine-6-phosphate deacetylase (EC 3.5.1.25) [Lactococcus lactis súbsp. lactis]
SeqID 701	SA-1601.1	Contig139 (88835-89278 p)	55	Identities = 63/146 (43%), Positives = 87/146 (59%), Gaps = 1/146 (0%) sp[034589 FLAW_BACSU PROBABLE FLAVODOXIN 2 pir [E69866 flavodoxin homolog ykuP[similarity] Bacillus subtilis emb[CAA10879.1] (AJ222587) YkuP protein [Bacillus subtilis] emb[CAB13290.1] (Z99111) similar to sulfite reductase [Bacillus subtilis]



Identities = 123/343 (35%), Positives = 175/343 (50%), Gaps = 26/343 (7%) pir  T35340 probable adenosine deaminase - Streptomyces coelicolor emb CAB42949.1  (AL049863) putative adenosine deaminase [Streptomyces coelicolor A3(2)] Length = 359	Identities = 144/306 (47%), Positives = 209/306 (68%), Gaps = 4/306 (1%) pir  F6999 conserved hypothetical protein ytql - Bacillus subtilis gb AAC00337.1  (AF008220) Ytql [Bacillus subtilis] emb CAB14885.1  (299118) similar to hypothetical proteins [Bacillus subtilis] Length = 313	Identities = 67/89 (75%), Positives = 76/89 (85%) spl031587 R14B_BACSU 30S RIBOSOMAL PROTEIN S14-2 pir  F69835 ribosomal protein S14 homolog yhzA - Bacillus subtilis emb CAB12716.1  (299108) similar to ribosomal protein S14 [Bacillus subtilis] Length = 89	Identities = 139/339 (41%), Positives = 201/339 (59%), Gaps = 4/339 (1%) ref NP_047071.1  L4171.5 [Leishmania major] pir  T02833 threonine aldolase L4171.5 [imported] - Leishmania major (strain Friedlin) gb AAC24656.1  (AE001274) L4171.5 [Leishmania major]	Identities = 244/343 (71%), Positives = 280/343 (81%), Gaps = 7/343 (2%) gb AAK04393.1 AE006266_12 (AE006266) Osialoglycoprotein endopeptidase [Lactococcus lactis subsp. lactis] Length = 346	Identities = 44/141 (31%), Positives = 73/141 (51%), Gaps = 8/141 (5%) pir  B70351 ribosomal-protein-alanine acetyltransferase - Aquifex aeolicus gb AAC06803.1  (AE000696) ribosomal-protein-alanine acetyltransferase [Aquifex aeolicus]
51	99	92	89	18	48
Contig139 (87754-88776 p)	Contig139 (86553-87488 m)	Contig90 (12596-12865 m)	Contig90 (11209-12234 p)	Contig90 (10079-11089 p)	Contig90 (9557-10003 p)
SA-1602.1	SA-1603.2	SA-1604.1	SA-1605.2	SA-1606.2	SA-1607.1
SeqID 702	SeqID 703	SeqID 704	SeqID 705	SeqID 706	SeqID 707



SeqID 708	SA-1608.1	Contig90 (8905-9594 p)	45	Identities = 94/182 (51%), Positives = 127/182 (69%), Gaps = 6/182 (3%) emb CAA76861.1  (Y17797) hypothetical protein [Enterococcus faecalis] Length = 204
SeqID 709	SA-1609.1	Contig90 (8493-8723 m)	55	Identities = 30/67 (44%), Positives = 45/67 (66%), Gaps = 7/67 (10%) dbj BAB06380.1  (AP001516) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 69
SeqID 710	SA-161.2	Contig136 (90788-91228 m)	47	Identities = 37/142 (26%), Positives = 71/142 (49%), Gaps = 7/142 (4%) pirjB70057 transcription regulator MarR family homolog ywhA - Bacillus subtilis emb CAB02511.1  (280360) Unknown, similar to regulatory proteins [Bacillus subtilis] emb CAB15782.1  (299123) similar to transcriptional regulator (MarR family) [Bacillus subtilis] Length = 139
SeqID 711	SA-1610.1	Contig90 (6760-8439 m)	77	Identities = 351/550 (63%), Positives = 442/550 (79%) dbj BAB06381.1  (AP001516) unknown conserved protein [Bacillus halodurans] Length = 555
SeqID 712	SA-1611.1	Contig90 (6089-6598 p)	38	Identities = 45/119 (37%), Positives = 67/119 (55%), Gaps = 4/119 (3%) emb[CAA66624.1] (X97985) ORF1 [Staphylococcus aureus] Length = 255
SeqID 713	SA-1612.1	Contig90 (4595-5941 p)	88	Pos (U6 alaci
SeqID 714	SA-1613.1	Contig90 (4190-4561 p)	69	Identities = 59/123 (47%), Positives = 89/123 (71%), Gaps = 5/123 (4%) splP19083 GLNR_BACCE REGULATORY PROTEIN GLNR pir JJU0076 hypothetical 15K protein (glnA 5 region) - Bacillus cereus dbj BAA00402.1  (D00513) ORF129 [Bacillus cereus]
SeqID 715	SA-1614.1	Contig90 (3628-4110 p)	33	Identities = 35/146 (23%), Positives = 64/146 (42%), Gaps = 7/146 (4%) dbjjBAB04661.1  (AP001510) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 360

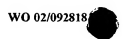


SeqID 716	SA-1615.1	Contig90 (2945-3355 m)	. 29	Identities = 61/134 (45%), Positives = 78/134 (57%), Gaps = 5/134 (3%) ref[NP_009939.1  Ycr013cp [Saccharomyces cerevisiae] sp[P25614 YCQ3_YEAST VERY HYPOTHETICAL 22.8 KD PROTEIN IN PGK1-POL4 INTERGENIC REGION pir  S19423 probable membrane protein YCR013c - yeast (Saccharomyces cerevisiae) emb[CAA42330.1  (X59720) YCR013c, len:215 [Saccharomyces cerevisiae] Length = 215
SeqID 717	SA-1616.1	Contig90 (2112-3308 p)	75	Identities = 254/398 (63%), Positives = 303/398 (75%), Gaps = 2/398 (0%) splQ9Z5C4 PGK_STAAU PHOSPHOGLYCERATE KINASE emb CAB38646.1  (AJ133520) phosphoglycerate kinase [Staphylococcus aureus]
SeqID 718	SA-1617.2	Contig90 (1108-1977 p)	81	Identities = 182/291 (62%), Positives = 237/291 (80%), Gaps = 7/291 (2%) emb CAA73175.1  (Y12602) acid phosphatase [Streptococcus equisimilis]
SeqID 719	SA-1619.1	Contig108 (18754-19293 m)	62	Identities = 89/146 (60%), Positives = 113/146 (76%), Gaps = 1/146 (0%) gb AAF08325.1 U78969_2 (U78969) FlaR [Streptococcus pyogenes]
SeqID 720	SA-162.1	Contig136 (88924-90741 m)	52	Identities = 216/543 (39%), Positives = 322/543 (58%), Gaps = 25/543 (4%) emb CAB69751.1  (AL137187) putative ABC transporter [Streptomyces coelicolor A3(2)] Length = 577
SeqID 721	SA-1620.1	Contig108 (17468-18553 m)	43	Identities = 94/307 (30%), Positives = 157/307 (50%), Gaps = 21/307 (6%) pir][G72378 glycerol dehydrogenase - Thermotoga maritima (strain MSB8) gblAAD35508.1 AE001721_7 (AE001721) glycerol dehydrogenase [Thermotoga maritima] Length = 364
SeqID 722	SA-1621.1	Contig108 (16596-17303 p)	23	Identities = 38/98 (38%), Positives = 59/98 (59%) emb[CAC01354.1  (AL390975) putative integral membrane protein [Streptomyces coelicolor A3(2)] Length = 198



SeqID 723	SA-1622.1	Contig108 (14405-16249 p)	69	Identities = 301/610 (49%), Positives = 437/610 (71%), Gaps = 9/610 (1%) dbjjBAB05348.1l (AP001512) unknown conserved protein [Bacillus halodurans]
SeqID 724	SA-1623.1	Contig108 (12108-14360 p)	, 89	dentities = 397/751 (52%), Positives = 519/751 (68%), Gaps = 14/751 (1%) dbj BAB04157.1  (AP001508) homosystein methyl transferase [Bacillus halodurans] Length = 756
SeqID 725	SA-1624.1	Contig108 (11622-11753 p)	No Hits found	
SeqID 726	SA-1625.2		09	Identities = 86/225 (38%), Positives = 142/225 (62%), Gaps = 1/225 (0%) gb AAK05711.1 AE006391_10 (AE006391) amino acid permease [Lactococcus lactis subsp. lactis]  235
SeqID 727	SA-1626.1	Contig87 (8382-8996 m)	No Hits found	
SeqID 728	SA-1627.1	Contig87 (7700-8284 m)	No Hits found	
SeqID 729	SA-1628.1	Contig87 (7137-7703 m)	No Hits found	
SeqID 730	SA-1629.1	Contig87 (4483-7137 m)	74	Identities = 544/880 (61%), Positives = 676/880 (76%), Gaps = 11/880 (1%) pir  SYBSVS valine-tRNA ligase (EC 6.1.1.9) - Bacillus stearothermophilus Length = 880
SeqID 731	SA-163.1	Contig136 (87180-88934 m)	55	Identities = 228/526 (43%), Positives = 331/526 (62%), Gaps = 2/526 (0%) emb CAB69752.1  (AL137187) putative ABC transporter [Streptomyces coelicolor A3(2)]
SeqID 732	SA-1630.1	Contig87 (3318-4247 p)	34	Identities = 57/231 (24%), Positives = 109/231 (46%), Gaps = 10/231 (4%) pir  T35598 hypothetical protein SC6G9.01c - Streptomyces coelicolor (fragment) emb CAB45592.1  (AL079356) hypothetical protein [Streptomyces coelicolor] Length = 409
SeqID 733	SA-1632.1	Contig87 (1942-2862 m)	43	Identities = 100/306 (32%), Positives = 152/306 (48%), Gaps = 3/306 (0%) pir  T50594 probable oxidoreductase [imported] - Streptomyces coelicolor emb CAB61731.1  (AL133220) putative oxidoreductase. [Streptomyces coelicolor A3(2)]  351

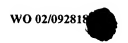
129



	90	n := 'r w	ω		00
Identities = 70/284 (24%), Positives = 135/284 (46%), Gaps = 8/284 (2%) ref[NP_053049.1  hypothetical protein [Plasmid pNZ4000] gb AAD40365.1  (AF036485) hypothetical protein [Plasmid pNZ4000]	Identities = 38/107 (35%), Positives = 59/107 (54%), Gaps = 6/107 (5%) splP37507 YYAQ_BACSU HYPOTHETICAL 13.9 KD PROTEIN IN COTF-TETB INTERGENIC REGION pir  S66000 yyaQ protein - Bacillus şubtilis dbj BAA05206.1  (D26185) unknown [Bacillus subtilis] emb CAB16112.1  (299124) yyaQ [Bacillus subtilis]	Identities = 234/384 (60%), Positives = 300/384 (77%), Gaps = 2/384 (0%) splP42602 YGJU_ECOLI HYPOTHETICAL SYMPORTER YGJU pir  F65097 hypothetical 43.5 kD protein in ebgC-exuT intergenic region - Escherichia coli (strain K-12) gb AAA57891.1  (U18997) ORF_0414; Geneplot suggests frameshift near start but none found [Escherichia coli] gb AAC76124.1  (AE000391) putative transport protein [Escherichia coli K12] Length = 414	Identities = 109/182 (59%), Positives = 141/182 (76%)   emb CAB59830.1  (AJ012388) hypothetical protein [Lactococcus lactis]   Length = 182	Identities = 67/182 (36%), Positives = 111/182 (60%), Gaps = 4/182 (2%) dbj BAB05604.1  (AP001513) unknown conserved protein [Bacillus halodurans]	Identities = 150/481 (31%), Positives = 247/481 (51%), Gaps = 15/481 (3%) pir  H69858 cation ABC transporter (ATP-binding protein) homolog ykoD - Bacillus subtilis emb CAA05601.1  (AJ002571) YkoD [Bacillus subtilis] emb CAB13179.1  (Z99110) similar to cation ABC transporter (ATP-binding protein) [Bacillus subtilis] Length = 490
44	23	7.3	61	51	44
Contig87 (879-1781 m)	Contig87 (3-611 m)	Contig124 (24911-26119 m)	Contig124 (24194-24742 m)	Contig136 (86446-87072 m)	Contig124 (22501-24177 m)
SA-1634.1	SA-1635.1	SA-1636.2	SA-1638.1	SA-164.1	SA-1640.2
SeqID 734	SeqID 735	SeqID 736	SeqID 737	SeqID 738	SeqID 739



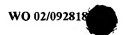
SeqID 740	SA-1641.2	Contig124 (21678-22508 m)	25	Identities = 46/134 (34%), Positives = 74/134 (54%), Gaps = 3/134 (2%) pir[JF75200 hypothetical protein PAB2261 - Pyrococcus abyssi (strain Orsay) emb[CAB49053.1] (AJ248283) hypothetical protein [Pyrococcus abyssi]
SeqID 741	SA-1642.1	Contig124 (20972-21643 p)	49	dentities = 70/183 (38%), Positives = 113/183 (61%), Gaps = 2/183 (1%) gb AAC46144.1  (AF001974) putative TrkA [Thermoanaerobacter ethanolicus] Length = 195
SeqID 742	SA-1643.1	Contig124 (19568-20959 p)	55	Identities = 170/461 (36%), Positives = 263/461 (56%), Gaps = 28/461 (6%) pir  G53610 ntpJ protein - Enterococcus hirae Length = 448
SeqID 743	SA-1644.1	Contig124 (18852-19565 p)	02	Identities = 132/240 (55%), Positives = 171/240 (71%), Gaps = 4/240 (1%) splP25813 GIDB_BACSU GLUCOSE INHIBITED DIVISION PROTEIN B pir  BWBSGB glucose-inhibited division protein gidB - Bacillus subtilis emb CAA44405.1  (X62539) homologous to E.coli gidB [Bacillus subtilis] dbj BAA05230.1  (D26185) unknown [Bacillus subtilis] emb CAB16137.1  (Z99124) glucose-inhibited division protein [Bacillus subtilis] Length = 239
SeqID 744	SA-1645.1	Contig124 (18170-18724 m)	78	Identities = 124/182 (68%), Positives = 152/182 (83%) gb AAG23700.1  (AF017421) LemA-like protein [Streptococcus gordonii] Length = 189
SeqID 745	SA-1646.1	Contig124 (17259-18149 m)	 <b></b>	Identities = 220/297 (74%), Positives = 261/297 (87%), Gaps = 1/297 (0%) splO30795 HTPX_STRGC PROBABLE PROTEASE HTPX HOMOLOG pir[ T48855 probable heat shock protein HtpX [imported] - Streptococcus gordonii gb AAB70525.1  (AF017421) putative heat shock protein HtpX [Streptococcus gordonii] Length = 297
SeqID 746	SA-1647.1	Contig124 (16621-17154 m)	70	Identities = 92/174 (52%), Positives = 133/174 (75%), Gaps = 3/174 (1%) gb AAG32547.1  (U12643) YlbN-like hypothetical protein [Streptococcus gordonii] Length = 176



SeqID 747	SA-1648.1	Contig124 (15697-16386 m)	91	Identities = 193/229 (84%), Positives = 211/229 (91%), Gaps = 1/229 (0%) gb AAC64935.1  (AF082668) CsrR [Streptococcus pyogenes] Length = 228
SeqID 748	SA-1649.2	Contig124 (14202-15707 m)	72	Identities = 248/501 (49%), Positives = 364/501 (72%), Gaps = 4/501 (0%) gb AAC64936.1  (AF082668) CsrS [Streptococcus pyogenes] Length = 500
SeqID 749	ŠA-165.1	Contig136 (85758-86465 m)	58	Identities = 84/226 (37%), Positives = 138/226 (60%), Gaps = 9/226 (3%) dbj BAB04091.1  (AP001508) two-component response regulator [Bacillus halodurans] Length = 222
SeqID 750	SA-1650.2	Contig134 (63460-63936 p)	41	Identities = 49/136 (36%), Positives = 69/136 (50%), Gaps = 8/136 (5%) pir[]G75571 MutT/nudix family protein - Deinococcus radiodurans (strain R1) gb[AAF09597.1]AE001864_3 (AE001864) MutT/nudix family protein [Deinococcus radiodurans] Length = 350
SeqID 751	SA-1651.1	Contig134 (64258-65064 p)	47	Identities = 70/254 (27%), Positives = 127/254 (49%), Gaps = 2/254 (0%) pir] G83412 hypothetical protein PA1860 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG05249.1 AE004612_3 (AE004612) hypothetical protein [Pseudomonas aeruginosa] Length = 275
SeqID 752	SA-1653.1	Contig134 (65459-65956 p)	. 84	Identities = 140/164 (85%), Positives = 157/164 (95%)   gb AAC38046.1  (AF000954) No definition line found   Streptococcus mutans  Length = 164
SeqID 753	SA-1654.1	Contig134 (65937-66335 p)	06	Identities = 109/133 (81%), Positives = 122/133 (90%), Gaps = 2/133 (1%) pir  A36933 diacylglycerol kinase homolog - Streptococcus mutans
SeqID 754	SA-1655.1	Contig134 (66332-67276 p)	83	Identities = 273/299 (91%), Positives = 290/299 (96%) gb AAD41632.1 AF072811_1 (AF072811) GTPase Era [Streptococcus pneumoniae] Length = 299
SeqID 755	SA-1656.1		No Hits found	
SedID 756	SA-1657.1	Contig134 (68290-68625 p)	No Hits found	



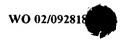
SeqID 757	SA-1658.1	Contig134 (68686-68898 p)	25	Identities = 31/87 (35%). Positives = 48/87 (54%), Gaps = 1/87 (1%) emb CAC03528 1  (AJ276410) BipY protein [Streptococcus pneumon ae] Length = 229
SeqID 758	SA-1659.1	Contig134 (69367-69447 p)	No Hits found	
SeqID 759	SA-166.1	Contig 136 (84529-85758 m)	38	Identities = 76/315 (24%). Positives = 149/315 (47%), Gaps = 33/315 (10%) gblAAF79919 1j (AF039082) putative histidine protein kinase [Lactococcus lactis]
SeqID 760	SA-1660.1	Contig134 (69841-71433 m)	10	Identities = 36/119 (30%), Positives = 57/119 (47%), Gaps = 15/119 (12%) pir  S74641 hypothetical protein sll1681 - Synechocystis sp. (strain PCC 6803) dbj BAA16793.1  (D90900) hypothetical protein [Synechocystis sp.]
SeqID 761	SA-1662.1	Contig134 (71635-72498 p)	48	Identities = 71/273 (26%), Positives = 140/273 (51%), Gaps = 16/273 (5%) splP49330JRGG_STRGC RGG PROTEIN pir]JA41898 positive regulator of glucosyltransferase and Spp phenotype rgg - Streptococcus gordonii gbJAAA26968.1 (M89776) rgg [Streptococcus gordonii] Length = 297
SeqID 762	SA-1663.1	Contig134 (72647-73468 p)	. 22	Identities = 182/271 (67%), Positives = 217/271 (79%) splP55045 FPG_STRMU FORMAMIDOPYRIMIDINE-DNA GLYCOSYLASE (FAPY-DNA GLYCOSYLASE) dbjlBAA05066.1  (D26071) formamidopyrimidine-DNA glycosylase [Streptococcus mutans] Length = 273
SeqID 763	SA-1665.1	Contig134 (73444-74052 p)	25	Identities = 80/189 (42%), Positives = 113/189 (59%), Gaps = 1/189 (0%) sp[O34932 YTAG_BACSU HYPOTHETICAL 22.0 KD PROTEIN IN GAPB-MUTM INTERGENIC REGION pir  A69988 conserved hypothetical protein ytaG - Bacillus subtilis gb AAC00353.1  (AF008220). YtaG [Bacillus subtilis] emb CAB14866.1  (Z99118) similar to hypothetical proteins [Bacillus subtilis]



SeqID 764	SA-1667.1	Contig134 (74176-74877 p)		Identities = 68/247 (27%), Positives = 126/247 (50%), Gaps = 17/247 (6%) pir  E69771 ABC transporter (ATP-binding protein) homolog ydbJ - Bacillus subtilis dbj BAA19286.1  (AB001488) PROBABLE TRANSPORT ATP BINDING PROTEIN. [Bacillus subtilis] emb CAB12256.1  (Z99106) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis]
SeqID 765	SA-1669.1	Contig134 (74855-76513 p)	No Hits found	
SeqID 766	SA-167.1	Contig136 (83535-84413 m)	74	Identities = 183/290 (63%), Positives = 223/290 (76%) gb[AAG02450.1 AF290097_1 (AF290097) mevalonate kinase [Streptococcus pyogenes]
SeqID 767	SA-1671.2	Contig134 (76757-77878 p)	. 22	Identities = 230/396 (58%), Positives = 315/396 (79%) emb[CAA07482.1  (AJ007367) multi-drug resistance efflux pump [Streptococcus pneumoniae] Length = 399
SeqID 768	SA-1673.2	Contig133 (6150-7757 m)		Identities = 413/535 (77%), Positives = 476/535 (88%), Gaps = 1/535 (0%) splQ59905IDEXB_STREQ GLUCAN 1,6-ALPHA-GLUCOSIDASE (DEXTRAN GLUCOSIDASE) (EXO-1,6-ALPHA-GLUCOSIDASE) (GLUCODEXTRANASE) pir  S39970 glucan 1,6-alpha-glucosidase (EC 3.2.1.70) - Streptococcus equisimilis emb CAA51348.1  (X72832) glucan 1,6-alpha-glucosidase [Streptococcus equisimilis] Length = 537
SeqID 769	SA-1674.1	Contig133 (5067-6062 m)	91	Identities = 267/331 (80%), Positives = 306/331 (91%) splP96995 GALE_STRMU UDP-GLUCOSE 4-EPIMERASE (GALACTOWALDENASE) (UDP-GALACTOSE 4-EPIMERASE EPIMERASE) (UDP-GALACTOSE 5-1.3.2) - Streptococcus mutans gb AAB49738.1  (U21942) UDP-galactose 4-epimerase (Streptococcus mutans) Length = 333



Identities = 71/223 (31%), Positives = 139/223 (61%), Gaps = 7/223 (3%) dbj BAB06470.1 (AP001516) two-component response regulator [Bacillus halodurans]	Identities = 132/461 (28%), Positives = 245/461 (52%), Gaps = 7/461 (1%) pir  A70009 two-component sensor histidine kinase homolog yufL - Bacillus subtilis emb CAB07946.1  (293937) unknown [Bacillus subtilis] emb CAB15141.1  (299120) similar to two-component sensor histidine kinase [YufM] [Bacillus subtilis]	Identities = 329/428 (76%), Positives = 375/428 (86%) 82 gb AAB18291.1  (U35658) L-malate permease [Streptococcus bovis] Length = 441	Identities = 36/70 (51%), Positives = 44/70 (62%) pirjlB72732 hypothetical protein APE0395 - Aeropyrum pernix (strain K1) dbjjBAA79350 1j (AP000059) 127aa long hypothetical protein [Aeropyrum pernix] Length = 127	Identities = 285/386 (73%), Positives = 332/386 (85%), Gaps = 1/386 (0%) gb AAB07709.1  (U35659) malic enzyme [Streptococcus bovis] Length = 389	Identities = 220/312 (70%), Positives = 264/312 (84%)   gb AAG02456.1 AF290099_2 (AF290099) mevalonate   diphosphate decarboxylase [Streptococcus pneumoniae]   Length = 317	39%), G =00629	transcriptional regulator CtsR [Lactococcus lactis subsp.
	Contig133 (2946-4364 p)	Contig133 (1347-2684 m)	Contig133 (628-1212 p)	Contig133 (159-1322 m)	Contig136 (82609-83553 m)	Contig96 (2340-2804 p)	
SA-1675.1	SA-1676.1	SA-1677.1	SA-1678.2	SA-1679.2	SA-168.1	SA-1680.2	
SeqID 770	SeqID 771	SeqID 772	SeqID 773	SeqID 774	SeqiD 775	SeqID 776	



SeqID 778	SA-1682.1	Contig96 (5468-5959 m)	. 09	Identities = 53/156 (33%). Positives = 84/156 (52%) gb AAF15587 1 AF187951_1 (AF187951) promotes resistance to glutamine synthetase inhibitors [Activation-tagging vector pSKi015] Length = 183
SeqID 779	SA-1683.1	Contig96 (5973-6614 m)	85	Identities = 179/213 (84%). Positives = 197/213 (92%)   gb[AAC97156 1] (U49397) unknown [Streptococcus pyogenes]   Length = 213
SeqID 780	SA-1685.1	Contig96 (6735-7712 m)	93	Identities = 282/325 (86%), Positives = 306/325 (93%) gb[AAC97155.1] (U49397) unknown [Streptococcus pyogenes] Length = 325
SeqID 781	SA-1686.1	Contig96 (7696-8571 m)	. 84	Identities = 224/271 (82%), Positives = 247/271 (90%) splQ9ZB45 HSLO_STRPY 33 KDA CHAPERONIN (HEAT SHOCK PROTEIN 33 HOMOLOG) (HSP33) gb AAC97154.1  (U49397) unknown [Streptococcus pyogenes] Length = 277
SeqID 782	SA-1687.1	Contig96 (8711-9967 m)	34	Identities = 64/350 (18%), Positives = 146/350 (41%), Gaps = 7/350 (2%) gb AAB93480.1 (AF019377) tellurite resistance protein [Rhodobacter sphaeroides] Length = 396
SeqiD 783	SA-1689.2	Contig96 (9964-10782 m)	37	Identities = 56/238 (23%), Positives = 105/238 (43%), Gaps = 25/238 (10%) emb CAB39037.2  (AL034559) hypothetical protein, PFC0960c [Plasmodium falciparum] Length = 1946
SeqID 784	SA-169.1	Contig136 (81624-82616 m)	70	Identities = 170/330 (51%), Positives = 233/330 (70%), Gaps = 1/330 (0%) gb AAG02457.1 AF290099_3 (AF290099) phosphomevalonate kinase [Streptococcus pneumoniae] Length = 336
SeqID 785	SA-1690.2	Contig96 (10906-11172 m)	55	Identities = 35/76 (46%), Positives = 49/76 (64%) gb AAK03132.1  (AE006146) unknown [Pasteurella multocida] Length = 757
SeqID 786	SA-1694.1	Contig83 (3183-4526 m)	37	Identities = 143/211 (67%), Positives = 170/211 (79%), Gaps = 3/211 (1%) gb AAD00288.1  (U78607) putative secreted protein [Streptococcus mutans]
SeqID 787	SA-1695.1	Contig83 (2091-3071 m)	78	Identities = 209/312 (66%), Positives = 266/312 (84%), Gaps = 3/312 (0%) gb AAA62181.1  (M92842) prs [Listeria monocytogenes] Length = 318



Contig83 (808-1986 m)  Contig124 (4176-4679 m)  Contig124 (4765-6096 m)  Contig137 (28295-29017 p)  Contig136 (80632-81627 m)  Contig124 (6106-6696 m)	Identities = 261/391 (66%), Positives = 323/391 (81%) gb AAF06954.1 AF146529_1 (AF146529) aromatic amino acid aminotransferase [Lactococcus lactis subsp. cremoris] Length = 391	Identities = 81/242 (33%), Positives = 124/242 (50%), Gaps = 3/242 (1%) dbj BAB05088.1  (AP001511) unknown conserved protein [Bacillus halodurans] Length = 254	Identities = 70/150 (46%), Positives = 100/150 (66%), Gaps = 2/150 (1%) gb AAK06154.1 AE006436_3 (AE006436) UNKNOWN PROTEIN [Lactococcus lactis subsp. lactis] Length = 164	Identities = 237/432 (54%), Positives = 315/432 (72%), Gaps = 3/432 (0%) sp P40778 MURC_BACSU UDP-N-ACETYLMURAMATEALANINE LIGASE (UDP-N-ACETYLMURAMOYL-L-ALANINE SYNTHETASE) pir [C69662 UDP-N-acetylmuramatealanine ligase (EC 6.3.2.8) murC - Bacillus subtilis gb AAC00294.1  (AF008220) putative UDP-N-acetylmuramate-alanine ligase [Bacillus subtilis] emb CAB14957.1  (299119) UDP-N-acetyl muramate-alanine ligase [Bacillus subtilis]	Identities = 82/270 (30%), Positives = 124/270 (45%), Gaps = 34/270 (12%) pir  S45091 hypothetical protein iota - Streptococcus pyogenes plasmid pDB101 pir  S68125 hypothetical protein iota - Streptococcus pyogenes plasmid pSM19035 emb CA47097.1  (X66468) orf iota [Streptococcus pyogenes] Length = 268	Identities = 127/331 (38%), Positives = 195/331 (58%), Gaps = 9/331 (2%) dbj[BAB07793.1  (AB037666) hypothetical protein [Streptomyces sp. CL190] Length = 363	ind   Identities = 260/678 (38%). Positives = 405/678 (59%). Gaps =	20/01/2 (20/01/2) (20/01/2) (20/01/2) (20/01/2 (20/01/2) (
Contig124 Contig137 (Contig136 (Contig124 Contig124 Contig124 Contig124 Contig124	74	47	49	02	6	28	No Hits found	.88
	SA-1696.1 Contig83 (808-1986	SA-1697.1 Contig83 (57-818 r	SA-1698.2 Contig124 (4176-467	SA-1699.1 Contig124 (4765-609	_			SA-1702.1   Contia124 (6888-998

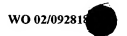
137

SeqID 796	SA-1704.1	Contig124 (10142-11452 m)	76	Identities = 377/436 (86%), Positives = 414/436 (94%) dbj BAA88823.1  (AB016077) phosphoglycerate dehydrogenase [Streptococcus mutans] Length = 436
SeqID 797	SA-1705.1	Contig124 (11500-12402 m)	63	Identities = 105/313 (33%), Positives = 191/313 (60%), Gaps = 17/313 (5%) splP06567 DNAL_BACSU PRIMOSOMAL PROTEIN DNA! pir  UQBS44 primosome component (helicase loader) dnal-Bacillus subtilis emb CAA28633.1  (X04963) ORF 311 (AA 1-311) [Bacillus subtilis] emb CAA99605.1  (Z75208) replication protein [Bacillus subtilis] gb AAC00359.1  (AF008220) Dnal [Bacillus subtilis] emb CAB14858.1  (Z99118) helicase loader [Bacillus subtilis]
SeqID 798	SA-1706.2	Contig124 (12399-13574 m)	. 25	Identities = 125/388 (32%), Positives = 207/388 (53%), Gaps = 14/388 (3%) gbjAAK04849.1 AE006308_9 (AE006308) replication protein DnaB [Lactococcus factis subsp. Length = 391
SeqID 799	SA-1707.2	Contig124 (13574-14053 m)	27	Identities = 100/150 (66%), Positives = 119/150 (78%), Gaps = 2/150 (1%) gb AAK04848.1 AE006308_8 (AE006308) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 148
SeqID 800	SA-171.1	Contig136 (80346-80564 m)	No Hits found	
SeqID 801	SA-1710.1	Contig98 (11651-12985 m)	No Hits found	
SeqID 802	SA-1711.1	Contig98 (10742-11626 p)	42	Identities = 97/251 (38%), Positives = 147/251 (57%), Gaps = 15/251 (5%) splP54604 YHCT_BACSU HYPOTHETICAL 33.7 KD PROTEIN IN CSPB-GLPP INTERGENIC REGION pir  H69823 conserved hypothetical protein yhcT - Bacillus subtilis emb CA465704.1  (X96983) hypothetical protein [Bacillus subtilis] emb CA812749.1  (Z99108) similar to hypothetical proteins [Bacillus subtilis]
SeqID 803	SA-1712.1	Contig98 (8377-10698 m)	12	Identities = 424/773 (54%), Positives = 555/773 (70%), Gaps = 47/773 (6%) gblAAF04735.1 AF101780_1 (AF101780) penicillinbinding protein 2a [Streptococcus pneumoniae] Length = 731



SeqID 804	SA-1714.1	Contig98 (7967-8140 m)	No Hits found	
	SA-1715.2	Contig98 (5728-6195 p)	49	dentities = 54/136 (39%), Positives = 77/136 (55%), Gaps = 19/136 (13%) pir  JQ0138 hypothetical 18.2K protein - Pseudomonas aeruginosa Length = 202
1	SA-1716.2	Contig98 (5724-7655 m)	22	Identities = 96/302 (31%), Positives = 143/302 (46%), Gaps = 37/302 (12%) gb AAB71985.1  (U73163) M-like protein [Streptococcus equi]
	SA-1718.2	Contig102 (7846-9225 p)	54	Identities = 147/473 (31%), Positives = 256/473 (54%), Gaps = 34/473 (7%) pir[ D69159 methyl coenzyme M reductase system, component A2 homolog - Methanobacterium thermoautotrophicum (strain Delta H) gb AAB84960.1  (AE000829) methyl coenzyme M reductase system, component A2 homolog [Methanothermobacter thermautotrophicus] Length = 480
	SA-1719.1	Contig102 (7122-7853 p)	36	Identities = 49/192 (25%), Positives = 89/192 (45%), Gaps = 6/192 (3%) pir  C69159 conserved hypothetical protein MTH453 - Methanobacterium thermoautotrophicum (strain Delta H) gb AAB84959.1  (AE000829) conserved protein [Methanobacterium thermoautotrophicum]
	SA-172.1	Contig136 (79446-80300 p)	48	Identities = 86/291 (29%), Positives = 151/291 (51%), Gaps = 28/291 (9%) dbj BAB10885.1  (AB010693)   gene_id:K21C13.21~pir  T04769~strong similarity to unknown protein [Arabidopsis thaliana]
П	SA-1720.1	Contig102 (6544-7140 p)	No Hits found	
	SA-1722.1	Contig102 (6129-6533 p)	42	Identities = 38/153 (24%), Positives = 68/153 (43%), Gaps = 1/153 (0%) emb CAC17502.1  (AL450432) conserved hypothetical protein [Streptomyces coelicolor] Length = 677
1	SA-1723.1	Contig 102 (5905-6087 p)	No Hits found	

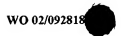
139



SeqID 813	SA-1724.1	Contig102 (5475-5918 p)	. 62	Identities = 77/221 (34%), Positives = 114/221 (50%), Gaps = 47/221 (21%) pir  T30345 irp5 protein - Yersinia enterocolitica emb CAA73130.1  (Y12527) Irp5 protein [Yersinia enterocolitica] Length = 525
SeqID 814	SA-1725.1	Contig102 (4539-5375 p)	59	Identities = 111/270 (41%), Positives = 165/270 (61%), Gaps = 2/270 (0%) sp P40871 DHBE_BACSU 2,3-DIHYDROXYBENZOATE-AMP LIGASE (DIHYDROXYBENZOIC ACID-ACTIVATING ENZYME) pir ID69615 2,3-dihydroxybenzoate[carrier protein] ligase (EC 6.2.1) dhbE-Bacillus subtilis gb AAC44632.1  (U26444) 2,3-dihydroxybenzoate-AMP ligase [Bacillus subtilis] emb CAB15188.1  (Z99120) 2,3-dihydroxybenzoate-AMP ligase (enterobactin synthetase component E) [Bacillus subtilis]
SeqID 815	SA-1726.1	Contig102 (3969-4343 p)	48	Identities = 43/113 (38%), Positives = 64/113 (56%), Gaps = 1/113 (0%) pir E71058 hypothetical protein PH1163 - Pyrococcus horikoshii dbj BAA30263.1  (AP000005) 148aa long hypothetical protein [Pyrococcus horikoshii]
SeqID 816	SA-1728.1	Contig102 (3469-3921 p)	. 29	Identities = 78/151 (51%), Positives = 108/151 (70%) dbj BAB05053.1  (AP001511) late competence operon required for DNA binding and uptake [Bacillus halodurans] Length = 188
SeqID 817	SA-1729.1	Contig102 (2390-3457 p)	89	Identities = 183/352 (51%), Positives = 258/352 (72%), Gaps = 3/352 (0%) emb CAA70068.1  (Y08842) aminopeptidase P [Lactococcus lactis] Length = 352
SeqID 818	SA-173.1	Contig136 (78431-79345 p)	51	Identities = 83/308 (26%), Positives = 160/308 (51%), Gaps = 12/308 (3%) pir[JF69795 conserved hypothetical protein yerQ - Bacillus subtilis emb[CAB12492.1] (Z99107) similar to hypothetical proteins [Bacillus subtilis] Length = 303
SeqID 819	SA-1730.1	Contig 102 (1574-2290 p)	No Hits found	



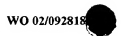
SA-1731.1	Contig102 (43-1572 p)	28	Identities = 198/481 (41%), Positives = 300/481 (62%), Gaps = 24/481 (4%) pir  F69763 multidrug resistance protein homolog ycnB - Bacillus subtilis dbj  BAA09016.1  (D50453) homologue of multidrug resistance protein B, EmrB, of E. coli [Bacillus subtilis] emb  CAB12192.1  (299106) similar to multidrug resistance protein [Bacillus subtilis] Length = 472
SA-1732.1 Contig130 (71572-71979 p)	979 p)	. 29	Identities = 73/135 (54%), Positives = 94/135 (69%), Gaps = 5/135 (3%) gb AAB99856.1  (AF043542) nucleoside diphosphate kinase [Gallus gallus]
1 Contig130 (	262 p) 358 p)	No Hits found No Hits found	
SA-1738.1 Contig130 (68962-69303 p)	(d E0	46	Identities = 44/104 (42%), Positives = 63/104 (60%), Gaps = -2/104 (1%) pir  F81982 hypothetical protein NMA0629 [imported] - Neisseria meningitidis (group A strain Z2491) emb CAB83918.1  (AL162753) hypothetical protein NMA0629 [Neisseria meningitidis] Length = 108
SA-1739.1 Contig130 (68096-68965 p)	5 p)	4	Identities = 71/224 (31%), Positives = 123/224 (54%) pir  G81982 hypothetical protein NMA0630 [imported] - Neisseria meningitidis (group A strain Z2491) emb CAB83919.1  (AL162753) hypothetical protein NMA0630 [Neisseria meningitidis] Length = 304
SA-174.1 Contig136 (77700-78344 p)	4 p)	28	Identities = 79/204 (38%), Positives = 132/204 (63%), Gaps = 4/204 (1%) spIP54176[HLY3_BACCE HEMOLYSIN III (HLY-III) pir  S59967 hemolysin III - Bacillus cereus emb CAA58877.1  (X84058) novel hemolytic factor [Bacillus cereus] Length = 219



SA-1740.2 Contig130 (63931-68094 p) 21 embj 90/765   SA-1741.2 Contig133 (58296-58787 p) 32 Streptot acress					
SA-1741.2 Contig133 (58296-58787 p) 32  SA-1742.1 Contig133 (58780-60048 p) 72  SA-1744.1 Contig133 (60591-60896 m) No Hits found SA-1746.1 Contig133 (60880-61281 m) No Hits found SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1748.1 Contig133 (62938-63327 m) No Hits found SA-1749.1 Contig133 (62938-63327 m) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig133 (63825-63741 m) No Hits found	SeqID 828	SA-1740.2	Contig130 (63931-68094 p)	21	CAI CAI
SA-1742.1 Contig133 (58780-60048 p) 72  SA-1744.1 Contig133 (60591-60896 m) No Hits found SA-1745.1 Contig133 (6059-62477 m) No Hits found SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1747.1 Contig133 (62938-63327 m) 30  SA-1748.1 Contig133 (63625-63741 m) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig133 (63625-64068 m) No Hits found	SeqID 829	SA-1741.2	Contig133 (58296-58787 p)	32	Identities = 35/109 (32%), Positives = 62/109 (56%), Gaps = 1/109 (0%) pir  T35305 probable acetyltransferase - Streptomyces coelicolor emb CAB51273.1  (AL096872) putative acetyltransferase [Streptomyces coelicolor A3(2)] Length = 169
SA-1744.1 Contig133 (60591-60896 m) No Hits found SA-1745.1 Contig133 (60880-61281 m) No Hits found SA-1745.1 Contig133 (61269-62477 m) No Hits found SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1748.1 Contig133 (62938-63327 m) 30 SA-1748.1 Contig133 (63625-63741 m) No Hits found SA-1750.1 Contig136 (77246-77695 p) No Hits found SA-1750.1 Contig133 (63625-64068 m) No Hits found SA-1750.1 Contig133 (63625-64068 m) No Hits found	SeqID 830	SA-1742.1	Contig133 (58780-60048 p)	72	Identities = 248/417 (59%), Positives = 315/417 (75%), Gaps = 4/417 (0%) pir  D69981 conserved hypothetical protein yrvN - Bacillus subtilis emb CAB14695.1  (299117) similar to hypothetical proteins [Bacillus subtilis] emb CAB14712.1  (299118) similar to hypothetical proteins [Bacillus subtilis]
SA-1745.1 Contig133 (60880-61281 m) No Hits found SA-1746.1 Contig133 (61269-62477 m) No Hits found SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1748.1 Contig133 (62938-63327 m) 30 SA-1749.1 Contig133 (63625-63741 m) No Hits found SA-1750.1 Contig136 (77246-77695 p) No Hits found SA-1750.1 Contig133 (63625-64068 m) No Hits found SA-1750.1 Contig133 (63625-64068 m) No Hits found	SeqID 831	SA-1744.1	Contig133 (60591-60896 m)	No Hits found	
SA-1746.1 Contig133 (61269-62477 m) No Hits found SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1748.1 Contig133 (62938-63327 m) 30 SA-1749.1 Contig133 (63625-63741 m) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig133 (63625-64068 m) No Hits found	SeqID 832	SA-1745.1	Contig133 (60880-61281 m)	No Hits found	
SA-1747.1 Contig133 (62479-62811 m) No Hits found SA-1748.1 Contig133 (62938-63327 m) 30 SA-1749.1 Contig133 (63625-63741 m) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig133 (63625-63741 m) No Hits found	SeqID 833	SA-1746.1	Contig133 (61269-62477 m)	No Hits found	
SA-1748.1 Contig133 (62938-63327 m) 30 SA-1749.1 Contig133 (63625-63741 m) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found SA-175.1 Contig136 (77246-77695 p) No Hits found	SeqID 834	SA-1747.1	Contig133 (62479-62811 m)	No Hits found	
SA-1749.1 Contig133 (63625-63741 m) SA-175.1 Contig136 (77246-77695 p) SA-1750.1 Contig133 (63820-64068 m)	SeqID 835	SA-1748.1	Contig133 (62938-63327 m)	30	Identities = 26/79 (32%), Positives = 47/79 (58%), Gaps = 1/79 (1%) pirl T13289 probable integrase - Streptococcus phage phi-01205 gb AAC79517.1  (U88974) ORF1 [Streptococcus thermophilus temperate bacteriophage O1205]
SA-175.1 Contig136 (77246-77695 p) SA-1750 1 Contig133 (63820-64068 m)	SeqID 836	SA-1749.1	Contig133 (63625-63741 m)	No Hits found	
SA-1750.1 Contio133 (63820-64068 m)	SeqID 837	SA-175.1	Contig 136 (77246-77695 p)	No Hits found	
'/	SeqID 838	SA-1750.1	Contig 133 (63820-64068 m)	No Hits found	



SA-1751.1	Contig133 (64228-64722 p)	27	Identities = 28/80 (35%), Positives = 49/80 (61%) pir  D69898 transcription regulator phage-related homolog yobD - Bacillus subtilis gb AAB84427.1  (AF027868) transcription regulator [Bacillus subtilis] emb CAB13777.1  (299114) similar to transcriptional regulator (phage-related) (Xre family) [Bacillus subtilis] Length = 112
SA-1753.2	Contig103 (9236-10060 p)	64	Identities = 138/268 (51%), Positives = 186/268 (68%), Gaps = 2/268 (0%) gb[AAK04909.1 AE006314_7 (AE006314) conserved hypothetical protein [Lactococcus lactis subsp. lactis]  Length = 269
SA-1754.1	Contig103 (8439-9236 p)	. 24	Identities = 91/264 (34%), Positives = 146/264 (54%), Gaps = 4/264 (1%) sp P75809 YBJ _ECOLI PROTEIN YBJ  Length = 271
	SA-1755.1 Contig103 (4809-8348 p)	61	Identities = 458/1193 (38%), Positives = 728/1193 (60%), Gaps = 27/1193 (2%) splP51834 SMC_BACSU CHROMOSOME PARTITION PROTEIN SMC pir  G69708 chromosome segregation SMC protein - Bacillus subtilis emb CAB13467.1  (Z99112) chromosome segregation SMC protein homolg [Bacillus subtilis] Length = 1186
	SA-1756.1 Contig103 (4115-4801 p)	63	Identities = 115/231 (49%), Positives = 155/231 (66%), Gaps = 1/231 (0%) splP51833 RNC_BACSU RIBONUCLEASE III (RNASE III) pirJB69693 ribonuclease III (EC 3.1.26.3) - Bacillus subtilis emb CAB13466.1  (299112) ribonuclease III [Bacillus subtilis] Length = 249
SA-1757.1	Contig103 (3571-3939 p)	54	Identities = 50/114 (43%), Positives = 72/114 (62%) emb CAC12789.1  (AJ279090) hypothetical protein [Staphylococcus carnosus]
SA-1758.1	Contig103 (2759-3568 p)	90	Identities = 223/269 (82%), Positives = 246/269 (90%) emb CAB65453.1  (AJ012051) VICX protein [Streptococcus pyogenes] Length = 270
SA-1759.1	Contig103 (1406-2755 p)	86	Identities = 335/443 (75%), Positives = 392/443 (87%) emb CAB65452.1  (AJ012051) VicK protein [Streptococcus pyogenes] Length = 452



SeqID 847	SA-176.1	Contig136 (75871-77154 p)	77	Identities = 266/421 (63%), Positives = 344/421 (81%), Gaps = 3/421 (0%) gb AAG02454.1 AF290098_2 (AF290098) HMG-CoA reductase [Streptococcus pneumoniae] Length = 424
SeqID 848	SA-1760.1	Contig103 (703-1413 p)	66	Identities = 205/237 (86%), Positives = 222/237 (93%) emb[CAB65451.1  (AJ012051) VicR protein [Streptococcus pyogenes] Length = 239
SeqID 849	SA-1761.1	Contig103 (15-215 p)	No Hits found	
SeqID 850	SA-1762.2	Contig118 (25352-25687 p)	57	Identities = 33/110 (30%), Positives = 65/110 (59%) gb AAC62417.1  (AF084104) hypothetical protein [Bacillus firmus] Length = 118
SeqID 851	SA-1763.1	Contig118 (24102-25268 p)	73	Identities = 227/390 (58%), Positives = 284/390 (72%), Gaps = 2/390 (0%) dbj BAB05375.1  (AP001512) chorismate synthase [Bacillus halodurans] Length = 390
SeqID 852	SA-1764.1	Contig118 (23034-24101 p)	55	Identities = 137/351 (39%), Positives = 199/351 (56%), Gaps = 4/351 (1%) splP73997IAROB_SYNY3 3-DEHYDROQUINATE SYNTHASE pir  S75507 3-dehydroquinate synthase - Synechocystis sp. (strain PCC 6803) dbj BAA18068.1  (D90911) 3-dehydroquinate synthase [Synechocystis sp.] Length = 361
SeqID 853	SA-1765.1	Contig118 (22263-22940 p)	53	Identities = 70/233 (30%), Positives = 127/233 (54%), Gaps = 12/233 (5%) splP35146 AROD_BACSU 3-DEHYDROQUINATE DEHYDRATASE (3-DEHYDROQUINASE) (TYPE I DHQASE) pir  S45563 3-dehydroquinate dehydratase (EC 4.2.1.10) aroC - Bacillus subtilis gb AA67501.1  (L09228) dehydroquinate dehydratase [Bacillus subtilis] emb CAB14240.1  (299116) 3-dehydroquinate dehydratase [Bacillus subtilis]



SeqID 854	SA-1766.1	Contig118 (21106-22263 p)	59	Identities = 161/396 (40%), Positives = 234/396 (58%), Gaps = 11/396 (2%) splP39587 YWBD_BACSU HYPOTHETICAL 44.4 KD PROTEIN IN EPR-GALK INTERGENIC REGION pir  S39674 ywbD protein - Bacillus subtilis emb CAA51575.1  (X73124) ipa-194 [Bacillus subtilis] emb CAB1562.1  (Z99123) alternate gene name: ipa-19d~similar to hypothetical proteins [Bacillus subtilis]
SeqID 855	SA-1767.1	Contig118 (18827-20971 m)	57	Identities = 298/682 (43%), Positives = 417/682 (60%), Gaps = 33/682 (4%) pir D69815 conserved hypothetical protein yfnl - Bacillus subtilis dbj BAA20118.1  (D86418) Yfnl [Bacillus subtilis] emb CAB12545.1  (299107) alternate gene name: yetP~similar to hypothetical proteins [Bacillus subtilis] Length = 653
SeqID 856	SA-1768.2	Contig118 (18227-18586 p)	16	Identities = 101/119 (84%), Positives = 110/119 (91%)     gb AAK05940.1 AE006414_6 (AE006414) 50S ribosomal protein     L20 [Lactococcus lactis subsp.   lactis]   Length = 119
SeqID 857	SA-1769.2	Contig118 (15777-15974 m)	. 44	Identities = 36/65 (55%), Positives = 49/65 (75%), Gaps = 2/65 (3%) gb AAK05803.1 AE006401_4 (AE006401) ferredoxin
SeqID 858	SA-177.1	Contig136 (74670-75869 p)	72	Identities = 261/385 (67%), Positives = 325/385 (83%)   gb AAG02448.1 AF290096
SeqID 859	SA-1770.1	Contig118 (15299-15790 p)	, 40	Identities = 44/135 (32%), Positives = 66/135 (48%), Gaps = 10/135 (7%) splP36920 EBSA_ENTFA PORE FORMING PROTEIN EBSA pir  A49939 ebsA protein - Enterococcus faecalis gb AAC36851.1  (L23802) pore-forming peptide [Enterococcus faecalis]

SeqID 860	SA-1771.1	Contig118 (14050-15270 p)	18	Identities = 274/406 (67%), Positives = 334/406 (81%), Gaps = 4/406 (0%) splP42020IPEPT_LACLC PEPTIDASE T (AMINOTRIPEPTIDASE) (TRIPEPTIDASE) gblAAA20627.1  (L27596) tripeptidase [Lactococcus lactis] Length = 413 Identities = 178/545 (32%), Positives = 302/545 (54%), Gaps = 26/545 (4%) pir  G69992 spore cortex protein homolog ytgP - 26/545 (4%) pir   G69992 spore cortex protein homolog ytgP - 26/545 (4%) pir   G69992 spore cortex protein homolog ytgP - 26/545 (4%) pir
SeqID 862	SA-1773.1	Contig 118 (10702-1296 m)	47	subtilis gp/AC00270.1  (AF008220) YtgP [Bacillus subtilis] emb CAB14983.1  (Z99119) similar to spore cortex protein [Bacillus subtilis] Length = 544  Identities = 153/468 (32%), Positives = 237/468 (49%), Gaps = 23/468 (4%) dbj BAB06290.1  (AP001515) UDP-N-acetylmuramoylalanyl-D-glutamyl-2, 6-diaminopimelate ligase [Bacillus halodurans] Length = 486
SeqID 863	SA-1774.2	Contig118 (9753-10547 m)	89	Identities = 141/259 (54%), Positives = 193/259 (74%) gb AAD33527.2 AF132117_4 (AF132117) FhuA (Staphylococcus aureus] gb AAF98153.1 AF251216_1 (AF251216) FhuC [Staphylococcus aureus] Length = 265
SeqID 864	SA-1775.2	C <sub>u</sub> ntig118 (8797-9729 m)	55	Identities = 95/301 (31%), Positives = 178/301 (58%), Gaps = 11/301 (3%) dbj BAB07609.1  (AP001520) ferrichrome ABC transporter (ferrichrome-binding protein) [Bacillus halodurans] Length = 308
SeqID 865	SA-1776.1	Contig118 (7756-8781 m)	56	Identities = 117/313 (37%), Positives = 195/313 (61%), Gaps = 3/313 (0%) gb AAD33526.3 AF132117_3 (AF132117) ferrichrome transport permease [Staphylococcus aureus] gb AAF98154.1 AF251216_2 (AF251216) FhuB [Staphylococcus aureus] aureus]
SeqID 866	SA-1778.1	Contig118 (6758-7759 m)	61	Identities = 122/334 (36%), Positives = 210/334 (62%), Gaps = 3/334 (0%) gb AAD33524.3 AF132117_1 (AF132117) ferrichrome transport permease [Staphylococcus aureus] gb AAF98155.1 AF251216_3 (AF251216) FhuG [Staphylococcus aureus] Length = 338



SeqID 867	SA-1779.2	Contig118 (6075-6728 p)	09	Identities = 89/233 (38%), Positives = 128/233 (54%), Gaps = 23/233 (9%) gb AAK05909 1 AE006411_4 (AE006411) UNKNOWN PROTEIN [Lactococcus lactis subsp lactis] Lengin = 265
SeqID 868	SA-178.1	Contig136 (73704-74543 m)	83	Identities = 215/280 (76%), Positives = 244/280 (86%), Gaps = 2/280 (0%) splP19368 TYSY_LACLA THYMIDYLATE SYNTHASE (TS) (TSASE) pir  A43797 thymidylate synthase (EC 2 1 1 45) - Lactococcus lactis subsp. lactis gb AAA25221.1  (M33770) thymidylate synthase (EC 2.1.1.45)   [Lactococcus lactis] Length = 279
SeqID 869	SA-1780.2	Contig138 (28940-29971 m)	53	Identities = 137/350 (39%), Positives = 211/350 (60%), Gaps = 3/350 (0%) emb CAB54584.1  (AJ006400) histidine kinase [Streptococcus pneumoniae] Length = 365
SeqID 870	SA-1781.1	Contig138 (30022-30762 m)	51	Identities = 73/239 (30%), Positives = 127/239 (52%), Gaps = 4/239 (1%) emb CAB69806.1  (AJ243712) YVFS protein [Bacillus cereus]
SeqID 871	SA-1783.1	Contig138 (30749-31645 m)	42	Identities = 83/216 (38%), Positives = 136/216 (62%), Gaps = 1/216 (0%) gb AAG21390.1 AF302051_4 (AF302051) ABC transporter ATP binding subunit [Bacillus licheniformis] Length = 299
SeqID 872	SA-1784.1	Contig138 (31774-33384 m)	74	Identities = 303/533 (56%), Positives = 401/533 (74%), Gaps = 15/533 (2%) gbjAAF34762.1jAF228345_1 (AF228345) unknown [Listeria monocytogenes]
SeqID 873	SA-1785.1	Contig138 (33573-34055 p)	68	Identities = 129/160 (80%), Positives = 149/160 (92%)   gb AAG28749.1 AF295118_1 (AF295118) LuxS [Streptococcus   pyogenes] Length = 170
SeqID 874	SA-1787.1	Contig138 (34280-35743 m)	30	Identities = 71/398 (17%), Positives = 152/398 (37%), Gaps = 16/398 (4%) gb[AAD16120.1] (AF094508) dentin phosphoryn [Homo sapiens] Length = 788

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SeqID 880	SA-1794.1	Contig86 (4134-4607 m)	57	Identities = 56/161 (34%), Positives = 92/161 (56%), Gaps = 8/161 (4%) sp P39337 YJGM_ECOLI HYPOTHETICAL 18.6 KDA   PROTEIN IN ARGI-VALS INTERGENIC REGION Length = 167
SeqID 881	SA-1795.1	Contig86 (2328-4067 m)	28	Identities = 227/565 (40%), Positives = 342/565 (60%), Gaps = 1/565 (0%) emb[CAB69751.1] (AL137187) putative ABC transporter [Streptomyces coelicolor A3(2)]
SeqID 882	SA-1796.1	Contig86 (554-2245 m)	65	Identities = 270/611 (44%), Positives = 391/611 (63%), Gaps = 31/611 (5%) emb CAB69752.1  (AL137187) putative ABC transporter [Streptomyces coelicolor A3(2)]
SeqID 883	SA-1797.1	Contig86 (4-513 p)	. 69	Identities = 78/167 (46%), Positives = 118/167 (69%), Gaps = 1/167 (0%) pir  A33141 hypothetical protein (gtfD 3 region) - Streptococcus mutans Length = 169
SeqID 884	SA-1798.3	Contig93 (7851-9347 p)	65	Identities = 250/500 (50%), Positives = 329/500 (65%), Gaps = 4/500 (0%) splP29851 MALQ_STRPN 4-ALPHA-GLUCANOTRANSFERASE (AMYLOMALTASE) (DISPROPORTIONATING ENZYME) (D-ENZYME) gb AAA26923.1  (J01796) amylomaltase [Streptococcus pneumoniae] Length = 505
SeqiD 885	SA-1799.3	Contig93 (9359-11623 p)	99	Identities = 297/776 (38%), Positives = 452/776 (57%), Gaps = 41/776 (5%) splP39123lPHSG_BACSU GLYCOGEN PHOSPHORYLASE pir  S36628 glycogen phosphorylase (EC 2.4.1.1) - Bacillus subtilis pir  S40052 glycogen phosphorylase (EC 2.4.1.1) glgP - Bacillus subtilis emb CAA81044.1  (225795) Glycogen Phosphorylase [Bacillus subtilis] emb CAB15072.1  (299119) glycogen phosphorylase [Bacillus subtilis] emb CAB15072.1  (299119) glycogen phosphorylase [Bacillus subtilis]

SeqID 886	SA-18.1	Contig137 (25957-28218 p)	38.	Identities = 178/715 (24%), Positives = 273/715 (37%), Gaps = 117/715 (16%) pir  G41662 130K surface exclusion protein Sec10 precursor - Enterococcus faecalis plasmid pCF10 gb AAA65847.1  (M64978) surface exclusion protein [Plasmid pCF10]
SeqID 887	SA-1801.2	Contig139 (45477-45692 p)	75	Identities = 28/76 (36%), Positives = 55/76 (71%), Gaps = 4/76 (5%) pir  E83139 exodeoxyribonuclease VII small subunit PA4042 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG07429.1 AE004821_2 (AE004821) exodeoxyribonuclease VII small subunit [Pseudomonas aeruginosa] Length = 80
SeqID 888	SA-1802.1	Contig139 (44159-45499 p)	63	Identities = 194/446 (43%), Positives = 284/446 (63%), Gaps = 10/446 (2%) splP54521 EX7L_BACSU PUTATIVE EXODEOXYRIBONUCLEASE LARGE SUBUNIT (EXONUCLEASE VII LARGE SUBUNIT) pirI G69960 exodeoxyribonuclease VII (large subunit) homolog yqiB - Bacillus subtilis dbj BAA12573.1  (D84432) YqiB  Bacillus subtilis] emb CAB14361.1  (299116) similar to exodeoxyribonuclease VII (large subunit)   Bacillus subtilis]
SeqID 889	SA-1803.1	Contig139 (43197-44033 p)	. 67	Identities = 149/277 (53%), Positives = 191/277 (68%) splP96051 YFOL_STRTR HYPOTHETICAL 29.9 KD PROTEIN IN FOLD-PBP2B INTERGENIC REGION (ORF1091) gb AAC44613.1  (U58210) orf1091 [Streptococcus thermophilus] Length = 278
SeqID 890	SA-1804.1	Contig139 (42346-43200 p)	83	Identities = 209/282 (74%), Positives = 248/282 (87%) sp[P96050 FOLD_STRTR FOLD BIFUNCTIONAL PROTEIN [INCLUDES: METHYLENETETRAHYDROFOLATE DEHYDROGENASE : METHENYLTETRAHYDROFOLATE CYCLOHYDROLASE



SeqID 891	SA-1806.1	Contig139 (40513-42207 p)	09	Identities = 238/548 (43%), Positives = 343/548 (62%), Gaps = 14/548 (2%) dbj BAB04825.1  (AP001510) phosphomannomutase [Bacillus halodurans] Length = 578
SeqID 892	SA-1807.2	Contig139 (39492-40226 p)	78	Identities = 153/239 (64%), Positives = 193/239 (80%) ref[NP_069514.1] glutamine ABC transporter, ATP-binding protein (glnQ) [Archaeoglobus fulgidus] pir  H69334 glutamine transport protein glnQ - Archaeoglobus fulgidus gb AAB90561.1  (AE001058) glutamine ABC transporter, ATP- binding protein (glnQ) [Archaeoglobus fulgidus] Length = 242
SeqID 893	SA-1808.2	Contig139 (38801-39499 p)	55	Identities = 92/209 (44%), Positives = 129/209 (61%), Gaps = 10/209 (4%) ref[NP_069070.1] glutamine ABC transporter, permease protein (glnP) [Archaeoglobus fulgidus] pir  H69278 glutamine ABC transporter, permease protein (glnP) homolog - Archaeoglobus fulgidus gb AAB91000.1  (AE001090) glutamine ABC transporter, permease protein (glnP) [Archaeoglobus fulgidus] Length = 224
SeqID 894	SA-1809.2	Contig89 (8855-9556 m)	89	Identities = 150/231 (64%), Positives = 178/231 (76%), Gaps = 1/231 (0%) splQ9ZHA7 DCOP_STRPN OROTIDINE 5 - PHOSPHATE DECARBOXYLASE (OMP DECARBOXYLASE) (OMPDCASE) gb AAC95452.1  (AF068902) orotidine-5 - decarboxylase PyrF [Streptococcus pneumoniae]
SeqID 895	SA-181.1	Contig136 (71686-72912 m)	85	Identities = 306/395 (77%), Positives = 357/395 (89%), Gaps = 1/395 (0%) gb AAF63738.1 AF236863_2 (AF236863) protease ClpX [Lactococcus lactis] Length = 411
SeqID 896	SA-1810.1	Contig89 (8213-8842 m)	84	Identities = 152/208 (73%), Positives = 180/208 (86%) gb AAC95453.1  (AF068902) orotate phosphoribosyltransferase PyrE [Streptococcus pneumoniae] Length = 210

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Identities = 176/422 (41%), Positives = 255/422 (59%), Gaps = 8/422 (1%) splO66990 PYRC_AQUAE DIHYDROOROTASE (DHOASE) pir  C70370 dihydroorotase - Aquifex aeolicus gb AAC06948.1  (AE000708) dihydroorotase [Aquifex aeolicus] Length = 422	Identities = 197/303 (65%), Positives = 250/303 (82%) gb AAF72727.1 (AF264709) aspartate transcarbamoylase [Enterococcus faecalis] Length = 308	Identities = 242/355 (68%), Positives = 305/355 (85%) emb CAB89872.1  (AJ132624) carbamoyl phosphate synthetase small subunit [Lactococcus lactis] Length = 357	Identities = 771/1062 (72%), Positives = 900/1062 (84%), Gaps = 5/1062 (0%) emb CAA03928.1  (AJ000109) carbamoylphosphate synthetase [Lactococcus lactis]		Identities = 39/130 (30%), Positives = 59/130 (45%), Gaps = 26/130 (20%) pirl T38271 conserved hypothetical protein SPAC23C4.16c - fission yeast (Schizosaccharomyces pombe) emb CAB16887.1  (299753) conserved hypothetical protein [Schizosaccharomyces pombe] Length = 424	Identities = 166/266 (62%), Positives = 207/266 (77%), Gaps = 2/266 (0%) gblAAK05380.1/AE006360_9 (AE006360) glutamate racemase (EC 5.1.1.3) [Lactococcus lactis subsp. lactis]	Identities = 96/189 (50%), Positives = 130/189 (67%), Gaps = 1/189 (0%) pir  C69986 conserved hypothetical protein ysnA - Bacillus subtilis emb CAA99555.1  (Z75208) hypothetical protein [Bacillus subtilis] emb CAB14796.1  (Z99118) similar to hypothetical proteins [Bacillus subtilis]
29	80	82	84	No Hits found	22	89	38
Contig89 (6909-8201 m)'	Contig89 (5821-6744 m)	Contig89 (4731-5807 m)	Contig89 (1518-4700 m)	Contig89 (1068-1391 m)	Contig89 (289-1071 m)	Contig91 (6891-7685 p)	Contig91 (7682-8656 p)
SA-1812.1	SA-1813.1	SA-1814.1	SA-1815.1	SA-1816.1	SA-1817.2	SA-1818.2	SA-1819.1
SeqID 897	SeqID 898	SeqID 899	SeqID 900	SeqID 901	SeqID 902	SeqID 903	SeqID 904



				Identities = 43/75 (57%), Positives = 57/75 (75%), Gaps = 1/75
SeqID 913	SA-1827.1	Contig91 (12363-12614 p)	99	(1%) pir[ G72251 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb[AAD36530.1 AE001797_10 (AE001797) conserved hypothetical protein [Thermotoga maritima] Length = 81
SeqID 914	SA-1828.1	Contig91 (12762-12881 p)	No Hits found	
SeqID 915	SA-183.1	Contig136 (70469-71098 p)	36	Identities = 44/172 (25%), Positives = 80/172 (45%), Gaps = 27/172 (15%) gb AAG09969.1 AF248037_4 (AF248037) unknown [Streptococcus agalactiae] Length = 174
SeqID 916	SA-1830.1	Contig91 (12638-14077 m)	53	Identities = 165/480 (34%), Positives = 261/480 (54%), Gaps = 10/480 (2%) ref[NP_069673.1] TRK potassium uptake system protein (trkH) [Archaeoglobus fulgidus] pir[JG69354 TRK potassium uptake system protein (trkH) homolog - Archaeoglobus fulgidus gb AAB90400.1] (AE001046) TRK potassium uptake system protein (trkH) [Archaeoglobus fulgidus] Length = 478
SeqID 917	SA-1832.1	Contig91 (14082-15431 m)	52	Identities = 136/446 (30%), Positives = 238/446 (52%), Gaps = 12/446 (2%) ref[NP_069672.1] TRK potassium uptake system protein (trkA-2) [Archaeoglobus fulgidus] sp[O29420]TRKA_ARCFU TRK SYSTEM POTASSIUM UPTAKE PROTEIN TRKA HOMOLOG pir ]F69354 TRK potassium uptake system protein (trkA-2) homolog - Archaeoglobus fulgidus gb[AAB90401.1] (AE001046) TRK potassium uptake system protein (trkA-2) [Archaeoglobus fulgidus] Length = 436
SeqID 918	SA-1834.1	Contig85 (10632-11486 p)	90	Identities = 91/289 (31%), Positives = 146/289 (50%), Gaps = 14/289 (4%) splP12055 STR_STAAU STREPTOMYCIN RESISTANCE PROTEIN pir  S00938 str protein - Staphylococcus aureus plasmid pS194 emb CAA29839.1  (X06627) ORF (str) [Staphylococcus aureus] Length = 282
SeqID 919	SA-1835.1	Contig85 (9974-10453 p)	No Hits found	
SeqID 920	SA-1836.1	Contig85 (8970-9626 p)	No Hits found	



/es = 57/103 (55%), Gaps = netical protein PAB0331 - amb CAB49414.1  (AJ248284) abyssi] Length = 114	iives = 54/110 (49%), Gaps = putative N-acetyltransferase b AAF22299.1 AF185571_1 ansferase Camello 2 [Homo Length = 227	/es = 94/188 (49%), Gaps = 1 protein yxlG - Bacillus subtilis pothetical [Bacillus subtilis] Bacillus subtilis] Length	ves = 176/301 (57%), Gaps = ACSU HYPOTHETICAL ABC PROTEIN 2 IN GLPD-CSPB 8822 ABC transporter (ATP-1 - Bacillus subtilis) lar to ABC transporter (ATP-1 subtilis) Length = 305	/es = 71/134 (52%), Gaps = 3767) ORF6 [Bacillus pumilus] 211	es = 105/186 (55%), Gaps = netical protein 2 - Bacillus   (221972) ORF2 [Bacillus Length = 216
Identities = 31/103 (30%), Positives = 57/103 (55%), Gaps = 4/103 (3%) pir  G75166 hypothetical protein PAB0331 - Pyrococcus abyssi (strain Orsay) emb CAB49414.1  (AJ248284) hypothetical protein [Pyrococcus abyssi]	Identities = 32/110 (29%), Positives = 54/110 (49%), Gaps = 4/110 (3%) rèfINP_057431.1  putative N-acetyltransferase Camello 2 [Homo sapiens] gblAAF22299.1 AF185571_1 (AF185571) putative N-acetyltransferase Camello 2 [Homo sapiens]	Identities = 42/188 (22%), Positives = 94/188 (49%), Gaps = 4/188 (2%) pir  B70082 hypothetical protein yxlG - Bacillus subtilis dbj BAA11738.1  (D83026) hypothetical [Bacillus subtilis] emb CAB15891.1  (299123) yxlG [Bacillus subtilis] Length = 259	Identities = 105/301 (34%), Positives = 176/301 (57%), Gaps = 11/301 (3%) splP54592 YHCH_BACSU HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN 2 IN GLPD-CSPB INTERGENIC REGION pir  D69822 ABC transporter (ATP-binding protein) homolog yhcH - Bacillus subtilis emb CAA65691.1  (X96983) hypothetical protein [Bacillus subtilis] emb CAB12736.1  (Z99108) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis] Length = 305	Identities = 39/134 (29%), Positives = 71/134 (52%), Gaps = 16/134 (11%) gb AAB71491.1  (U53767) ORF6 [Bacillus pumilus] Length = 211	Identities = 50/186 (26%), Positives = 105/186 (55%), Gaps 5/186 (2%) pir  S32217 hypothetical protein 2 - Bacillus megaterium emb CAA79986.1  (Z21972) ORF2 (Bacillus megaterium)
45	26	35	89	19	50
Contig85 (8753-9031 p)	Contig85 (7686-8174 p)	Contig85 (6726-7517 p)	Contig85 (5857-6759 p)	Contig85 (4576-5652 p)	Contig85 (3704-4327 p)
SA-1837.1	SA-1838.1	SA-1839.1	SA-1840.1	SA-1841.1	SA-1842.1
SeqID 921	SeqID 922	SeqID 923	SeqID 924	SeqID 925	SeqID 926



SeqID 927	SA-1843.1	Contig85 (3203-3499 m)	58	Identities = 53/96 (55%), Positives = 70/96 (72%) pir[ C70033 hypothetical protein yvdC - Bacillus subtilis emb CAB15470.1  (Z99121) yvdC [Bacillus subtilis] Length = 106
SeqID 928	SA-1844.1	Contig85 (2626-3186 m)	62	dentities = 83/185 (44%), Positives = 117/185 (52%) dbj BAB06803.1  (AP001517) unknown conserved protein [Bacillus halodurans] Length = 187
SeqID 929	SA-1845.1	Contig85 (1458-2333 m)	21	Identities = 40/132 (30%), Positives = 63/132 (47%), Gaps = 8/132 (6%) pir  E69787 hypothetical protein ydiL - Bacillus subtilis dbj  BAA19725.1  (D88802) transmembrane [Bacillus subtilis] emb  CAB12420.1  (299107) ydiL [Bacillus subtilis] Length = 244
SeqID 930	SA-1846.1	Contig85 (999-1301 p)	47	Identities = 38/92 (41%), Positives = 53/92 (57%), Gaps = 1/92 (1%) splP54510 YQHL_BACSU HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-SPOIIIAA INTERGENIC REGION pir  C69959 glpE protein homolog yqhL - Bacillus subtilis dbj BAA12549.1  (D84432) YqhL [Bacillus subtilis] emb CAB14385.1  (299116) similar to hypothetical proteins [Bacillus subtilis]
SeqID 931	SA-1847.1	Contig85 (59-826 p)	96	Identities = 252/255 (98%), Positives = 252/255 (98%) emb CAA51283.1  (X72754) cAMP factor [Streptococcus agalactiae] Length = 255
SeqID 932	SA-1849.1	Contig88 (13289-14137 p)	<b>5</b> 5	Identities = 94/276 (34%), Positives = 154/276 (55%), Gaps = 2/276 (0%) dbj BAB07346.1   (AP001519) unknown conserved protein [Bacillus halodurans] Length = 283
SeqID 933	SA-185.1	Contig 136 (69874-70248 m)	No Hits found	
SeqID 934	SA-1850.1	_	No Hits found	
SedID 935	SA-1851.1	Contig88 (12304-12837 p)	No Hits found	
SeqID 936	SA-1852.1	Contig88 (11759-12253 p)	. 61	Identities = 65/165 (39%), Positives = 102/165 (61%), Gaps = 5/165 (3%) gb[AAF66143.1] (U81488) TcsEorf2 [Lactococcus lactis subsp. cremoris]



	SA-1853.1	Contig88 (10706-11140 m)	37	Identities = 42/134 (31%), Positives = 55/134 (40%), Gaps = 18/134 (13%) dbjjBAA99921.1  (AP001306) contains similarity to cell wall-plasma membrane linker protein~gene_id:MKA23.5 [Arabidopsis thaliana] Length = 1480
	SA-1855.1	Contig88 (9606-11570 p)	99	Identities = 320/660 (48%), Positives = 439/660 (66%), Gaps = 46/660 (6%) dbj BAB04547.1  (AP001510) PTS system, fructosespecific enzyme II, BC component [Bacillus halodurans]  Length = 625
	SA-1858.1	Contig88 (8698-9609 p)	61	Identities = 146/303 (48%), Positives = 197/303 (64%) spl0317.14 K1PF_BACSU 1-PHOSPHOFRUCTOKINASE (FRUCTOSE 1-PHOSPHATE KINASE) pir  A69627 fructose 1-phosphate kinase fruB - Bacillus subtilis emb CAB13312.1  (299111) fructose-1-phosphate kinase [Bacillus subtilis] gb AAC24914.1  (AF012285) fructose-1-phosphate kinase [Bacillus subtilis]
ł	SA-1859.1	Contig88 (7958-8701 p)	53	Identities = 96/248 (38%), Positives = 148/248 (58%), Gaps = 1/248 (0%) dbj BAB04545.1 (AP001510) transcriptional repressor [Bacillus halodurans]
[ I	SA-186.1	Contig 136 (69066-69623 m)	No Hits found	
	SA-1860.1	Contig88 (6592-7827 p)	89	Identities = 216/410 (52%), Positives = 284/410 (68%) emb CAB89121.1  (AJ277485) beta-lactam resistance factor [Streptococcus pneumoniae]
,	SA-1861.1	Contig88 (4808-6448 p)	21	Identities = 93/183 (50%), Positives = 116/183 (62%), Gaps = 9/183 (4%) pir  C60328 hypothetical protein 2 (sr 5 region) - Streptococcus mutans (strain OMZ175, serotype f) Length = 179
	SA-1862.1	Contig88 (3679-4602 p)		Identities = 85/301 (28%), Positives = 150/301 (49%), Gaps = 7/301 (2%) pir  H75119 probable 2-dehydropantoate 2-reductase (EC 1.1.1.169) PAB0512 - Pyrococcus abyssi (strain Orsay) emb CAB49673.1  (AJ248285) PROBABLE 2-DEHYDROPANTOATE 2-REDUCTASE (EC 1.1.1.169) [Pyrococcus abyssi] Length = 300

SeqID 945	SA-1863.1	Contig88 (3120-3626 m)	No Hits found	
SeqID 946	SA-1866.2	Contig129 (15399-16343 m)	<b>5</b>	Identities = 111/300 (37%), Positives = 175/300 (58%), Gaps = 4/300 (1%) sp P37517 CCPB_BACSU CATABOLITE CONTROL PROTEIN B pir  S66011 transcription regulator homolog yyaG - Bacillus subtilis dbj BAA05217.1  (D26185) LACI family of transcriptional repressor (probable) [Bacillus subtilis] emb CAB16124.1  (299124) similar to transcriptional regulator (Lacl family) [Bacillus subtilis]
SeqID 947	SA-1867.1	Contig129 (14405-15340 p)	31	Identities = 63/218 (28%), Positives = 107/218 (48%), Gaps = 3/218 (1%) gb AAB17013.1  (L38252) esterase [Acinetobacter   woffii] Length = 303
SeqID 948	SA-1868.1	Contig129 (13444-14127 m)	06	Identities = 228/228 (100%), Positives = 228/228 (100%) emb[CAA72897.1] (Y12224) hypothetical protein [Streptococcus agalactiae] Length = 228
SeqID 949	SA-1869.1	Contig129 (12353-13390 m)	95	Identities = 343/345 (99%), Positives = 343/345 (99%) emb[CAA72898.1  (Y12224) hypothetical protein [Streptococcus agalactiae] Length = 345
SeqID 950	SA-1870.1	Contig129 (11665-12273 m)	. 26	Identities = 202/202 (100%), Positives = 202/202 (100%) splO54086 SODM_STRAG SUPEROXIDE DISMUTASE [MN-FE] Length = 202
SeqID 951	SA-1871.1	Contig129 (10476-11327 m)	63	Identities = 118/275 (42%), Positives = 183/275 (65%) dbj BAA11696.1  (D83026) LicT antiterminator [Bacilius subtilis] emb CAB15944.1  (Z99124) transcriptional antiterminator (BglG family) [Bacillus subtilis] Length = 277



	SA-1872.2	Contig129 (8615-10483 m)	55	Identities = 225/594 (37%), Postives = 351/594 (58%), Gaps = 11/594 (1%) spiP40739 PTBA_BACSU PTS SYSTEM, BETA-GLUCOSIDES.SPECIFIC IIABC COMPONENT (EIIABC-BGL) (BETA-GLUCOSIDES.PERMEASE IIABC COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, ABC COMPONENT) (EII-BGL) priji140406 beta-glucoside permease - Bacillus subtilis emb[CAA84286 1] (234526) beta-glucoside permease [Bacillus subtilis] Length = 609
SA-1	SA-1874.1	Contig115 (24509-25441 p)	***	Identities = 195/308 (63%), Positives = 245/308 (79%), Gaps = 4/308 (1%) spIP24137 OPPF_BACSU OLIGOPEPTIDE TRANSPORT ATP-BINDING PROTEIN OPPF pir  E38447 oligopeptide transport ATP-binding protein oppF - Bacillus subtilis gb AAA62692.1  (M57689) sporulation protein [Bacillus subtilis] Length = 308
SA-	SA-1875.1	Contig115 (23463-24509 p)	08	Identities = 230/342 (67%), Positives = 283/342 (82%), Gaps = 2/342 (0%) gb AAF73093.1 AF103793_4 (AF103793) ATPase OppD [Listeria monocytogenes] Length = 358
SA-	SA-1876.1	Contig115 (22419-23450 p)	63	Identities = 157/325 (48%), Positives = 219/325 (67%), Gaps = 4/325 (1%) gb AAF73092.1 AF103793_3 (AF103793) transmembrane protein OppC [Listeria monocytogenes] Length = 344
SA-	SA-1878.1	Contig115 (21495-22409 p)	69	Identities = 147/304 (48%), Positives = 221/304 (72%), Gaps = 1/304 (0%) gb AAF73091.1 AF103793_2 (AF103793) transmembrane protein OppB [Listeria monocytogenes] Length = 309
S, A	SA-1879.2	Contig115 (19721-21376 p)	74	Identities = 352/512 (68%), Positives = 416/512 (80%), Gaps = 2/512 (0%) gb AAD17886.1  (AF100456) hyaluronate-associated protein precursor [Streptococcus equi] Length = 522

Identities = 278/454 (61%), Positives = 351/454 (77%), Gaps = 3/454 (0%) pir  A69751 histidine permease homolog ybgF - Bacilius subtilis emb CAB12034.1  (Z99105) similar to histidine permease [Bacillus subtilis] dbj BAA33138.1  (AB006424) ybgF [Bacillus subtilis] Length = 470	Identities = 89/101 (88%), Positives = 94/101 (92%)   splQ9VVVV6 RL24_STRPN 50S RIBOSOMAL PROTEIN L24   gb AAD33267.1 AF126059_8 (AF126059) RpL24 [Streptococcus pneumoniae] gb AAD33276.1  (AF126060) RpL24   [Streptococcus pneumoniae] gb AAD33285.1  (AF126061)   RpL24 [Streptococcus pneumoniae]   Length = 101	Identities = 157/180 (87%), Positives = 172/180 (95%)   gb AAK06185.1 AE006437_15 (AE006437) 50S ribosomal   protein L5 [Lactococcus lactis subsp. lactis] Length   = 180	Identities = 100/132 (75%), Positives = 116/132 (87%) sp P12879 RS8_BACSU 30S RIBOSOMAL PROTEIN S8 (BS8) gb AAB06813.1  (L47971) ribosomal protein S8 [Bacillus subtilis] Length = 132	identities = 110/178 (61%), Positives = 134/178 (74%) spjP02391jRL6_BACST 50S RIBOSOMAL PROTEIN L6 (BL10) Length = 178	Identities = 86/120 (71%), Positives = 97/120 (80%), Gaps = 2/120 (1%) sp P46899 RL18_BACSU 50S RIBOSOMAL   PROTEIN L18_gb AAB06815.1  (L47971) ribosomal protein L18 [Bacillus subtilis] Length = 120	Identities = 119/158 (75%), Positives = 139/158 (87%) splP02357IRS5_BACST 30S RIBOSOMAL PROTEIN S5 (BS5) pirIJR3BS5F ribosomal protein S5 - Bacillus stearothermophilus abIAAA22699.11 (M57621) ribosomal protein S5 [Bacillus
74	78	06	98	72	92	18
Contig136 (67560-68948 m)	Contig132 (48543-48848 p)	Contig132 (48872-49414 p)	Contig132 (49772-50170 p)	Contig132 (50280-50816 p)	Contig132 (50917-51273 p)	Contig132 (51292-51786 p)
SA-188.1	SA-1881.2	SA-1882.2	SA-1883.1	SA-1884.1	SA-1885.1	SA-1886.1
SeqID 958	SeqID 959	SeqID 960	SeqID 961	SeqID.962	SeqID 963	SeqID 964



SA-1888.1	38.1	Contig132 (51801-51980 p)	29	Identities = 40/58 (68%), Positives = 46/58 (78%) sp O06444 \lambdaL30_STAAU 50S RIBOSOMAL PROTEIN L30 gb AAB54020.1  (U96620) ribosomal protein L30 {Staphylococcus aureus} Length = 59
SA-1891.1	91.1	Contig132 (52105-52545 p)	84	Identities = 116/146 (79%), Positives = 128/146 (87%) splO06445 RL15_STAAU 50S RIBOSOMAL PROTEIN L15 gb AAB54021.1  (U96620) ribosomal protein L15 {Staphylococcus aureus} Length = 146
SA-1892.1	92.1	Contig132 (52566-53870 p)	82	Identities = 292/433 (67%), Positives = 361/433 (82%), Gaps = 2/433 (0%) sp P27148 SECY_LACLA PREPROTEIN TRANSLOCASE SECY SUBUNIT pir  S17985 preprotein translocase secY - Lactococcus lactis subsp. lactis emb CAA41939.1  (X59250) SecY protein [Lactococcus lactis] prf  1715214A secY gene [Lactococcus lactis]
SA-1893.1	93.1	Contig132 (53965-54603 p)	7.7	Identities = 146/214 (68%), Positives = 170/214 (79%), Gaps = 6/214 (2%) splP27143jKAD_LACLA ADENYLATE KINASE (ATPAMP TRANSPHOSPHORYLASE) pirl S17987 adenylate kinase (EC 2.7.4.3) - Lactococcus lactis subsp. lactis pirl B44812 adenylate kinase (EC 2.7.4.3) - Lactococcus lactis emb CAA41940.1  (X59250) adenylate kinase [Lactococcus lactis] Length = 215

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Identities = 51/213 (23%). Positives = 89/213 (40%), Gaps = 32/213 (15%) pirt T19214 UDP-glucoseglycoprotein glucosyltransferase (EC 2 4 1 -) precursor F26H9.8 - Caenorhabditis elegans emb CAB03874.1  (281467) predicted using Genefinder~Similarity to Drosophila UDP-glucose glycoprotein glucosyltransferase (TR:Q09332), contains similarity to Pfam domain: PF01501 (Glycosyl transferase family 8), Score=-22.6, E-value=4.8e-05, N=1~cDNA EST yk250b10.3 co> emb CAB04207.1  (281516) predicted using Genefinder~Similarity to Drosophila UDP-glucose:glycoprotein glucosyltransferase (TR:Q09332), contains similarity to Pfam domain: PF01501 (Glycosyl transferase family 8), Score=-22.6, E-value=4.8e-05, N=1~cDNA EST yk250b10.3 co> Length = 1377	Identities = 324/433 (74%), Positives = 367/433 (83%) dbj BAB04344.1  (AP001509) adenylosuccinate lyase [Bacillus halodurans] Length = 433	Identities = 196/322 (60%), Positives = 254/322 (78%) spiO32055 RUVB_BACSU HOLLIDAY JUNCTION DNA HELICASE RUVB emb CAB75331.1  (Y15896) RuvB protein [Bacillus subtilis] Length = 334	Identities = 140/305 (45%), Positives = 191/305 (61%), Gaps = 6/305 (1%) splQ47690 MMUM_ECOLI HOMOCYSTEINE S-METHYLMETHIONINE:HOMOCYSTEINE METHYLMETHIONINE:HOMOCYSTEINE METHYLTRANSFERASE) pir  E64751 probable membrane protein yagD - Escherichia coli gb AAB08682.1  (U70214) similar to S. cerevisiae YLL062c [Escherichia coli] gb AAC73364.1  (AE000134) putative enzyme [Escherichia coli K12] Length
23	83	No Hits found	83
Contig132 (28026-29129 p)	Contig132 (29215-30513 p)	Contig132 (30700-31569 p) Contig132 (31858-32856 p)	Contig136 (66611-67555 m)

SA-1898.1

SeqID 972

SA-1899.1

SeqID 973

SA-190.1

SeqID 974

SA-1897.1

SeqID 971

SA-1894.2

SeqID 970



SA-1900.3		Contig132 (33008-33445 p)	61	Identities = 76/141 (53%), Positives = 100/141 (70%), Gaps = 4/141 (2%) gb AAK06300.1 AE006449_6 (AE006449) proteintyrosine phosphatase (EC 3.1.3.48) [Lactococcus lactis subsp. lactis] Length = 145
SA-1901.2 Conti	Contig	Contig106 (12011-12379 m)	. 62	Identities = 51/116 (43%), Positives = 79/116 (67%) dbj BAA83965.1  (AB024564) YHCF [Bacillus halodurans] dbj BAB04102.1  (AP001508) transcriptional regulator (GntR family) [Bacillus halodurans] Length = 126
SA-1902.2 Contig	Conti	Contig106 (12524-15628 p)	78	Identitles = 647/1036 (62%), Positives = 814/1036 (78%), Gaps = 4/1036 (0%) gb AAF98350.1  (AF280766) DNA polymerase III alpha subunit [Streptococcus pyogenes] Length = 1034
SA-1904.1 Contig	Contig	Contig106 (15709-16731 p)	88	Identities = 270/337 (80%), Positives = 302/337 (89%), Gaps = 1/337 (0%) gb AAF25803.1 AF172173_1 (AF172173) phosphofructokinase [Streptococcus thermophilus] Length = 339
SA-1906.1 Contig	Contig	Contig106 (16780-18282 p)	89	Identities = 413/500 (82%), Positives = 452/500 (89%) gb AAF25804.1 AF172173_2 (AF172173) pyruvate kinase [Streptococcus thermophilus] Length = 500
SA-1908.2 Contig106 (18	Contig	106 (18453-19010 p)	48	Identities = 62/185 (33%), Positives = 97/185 (51%), Gaps = 12/185 (6%) spIP72365ILEP_STAAU SIGNAL PEPTIDASE IB (SPASE IB) (LEADER PEPTIDASE IB) gb AAC44435.1  (U65000) type-1 signal peptidase SpsB [Staphylococcus aureus] Length = 191
SA-1909.2 Contig1	Contig1	Contig133 (13473-13898 m)	78	Identities = 92/141 (65%), Positives = 115/141 (81%) splP23494 LACA_LACLA GALACTOSE-6-PHOSPHATE ISOMERASE LACA SUBUNIT pir  A39778 galactose-6-phosphate isomerase (EC 5.3.1) component LacA - Lactococcus lactis gb AAA25168 1  (M65190) lacA [Lactococcus lactis] gb AAA25168 1  (M65190) lacA [Lactococcus lactis] gb AAA25177.1  (M60447) galactose 6-P isomerase [Lactococcus lactis]

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Identities = 437/589 (74%), Positives = 513/589 (86%), Gaps = 5/589 (0%) gb AAA68910.1  (L34677) Clp-like ATP-dependent protease binding subunit [Bos taurus]	Identities = 138/171 (80%), Positives = 157/171 (91%) sp P23495 LACB_LACLA GALACTOSE-6-PHOSPHATE ISOMERASE LACB SUBUNIT pir  B39778 galactose-6-phosphate isomerase (EC 5.3.1) component LacB - Lactococcus lactis gb AAA25169.1  (M65190) lacB [Lactococcus lactis] gb AAA25178.1  (M60447) galactose 6-P isomerase [Lactococcus lactis] Length = 171	Identities = 192/310 (61%), Positives = 236/310 (75%) sp P23391 LACC_LACLA TAGATOSE-6-PHOSPHATE KINASE (PHOSPHOTAGATOKINASE) pir  C39778 tagatose-6- phosphate kinase (EC 2.7.1) LacC - Lactococcus lactis gb AAA25170.1  (M65190) lacC [Lactococcus lactis] gb AAA25179.1  (M60447) tagatose 6-P kinase [Lactococcus lactis]	Identities = 253/325 (77%), Positives = 295/325 (89%) splP26593 LACD_LACLA TAGATOSE 1,6-DIPHOSPHATE ALDOLASE (TAGATOSE-BISPHOSPHATE ALDOLASE) (D-TAGATOSE-1,6-BISPHOSPHATE ALDOLASE) pir.  D39778 LacD tagatose-1,6-diphosphate aldolase - Lactococcus lactis gb AAA25171.1  (M65190) lacD [Lactococcus lactis] gb AAA25180.1  (M60447) tagatose 1,6-diP aldolase [Lactococcus lactis]	Identities = 173/298 (58%), Positives = 219/298 (73%) splP23496 LAXP_LACLA LACX PROTEIN, PLASMID pir  D23696 lacX protein - Lactococcus lactis gb AAA25184.1 (M60447) ORF [Lactococcus lactis] Length = 299
72	98	72	68	99
Contig136 (64409-66517 p)	Contig133 (12937-13452 m)	Contig133 (11994-12926 m)	Contig133 (11015-11992 m)	Contig133 (10065-10961 m)
SA-191.2	SA-1910.1	SA-1911.1	SA-1912.1	SA-1913.1
SeqID 982	SeqID 983	SeqID 984	SeqID 985	SeqID 986



0/278 (57%), Gaps = CINE RICH PROTEIN coccus equisimilis otein [Streptococcus ein [Strept	= 359/377 (94%) SUGAR-BINDING SMK pir[JE42400 ABC- n msmK [validated] - 177351) ATP-binding Length = 377	= 378/399 (94%) nosphopentomutase Length = 403	), Positives = 140/199 (69%), Gaps = 3 DEOC_MYCHO DEOXYRIBOSE-5 (PHOSPHODEOXYRIBOALDOLASE) E) pir  S42197 deoxyribose-phosphate   (SGC3)   Phosphate aldolase (EC 4.1.2.4) - (SGC3)   CSGC3) emp CA481646.1 (Z27121)   Coplasma hominis   Length = 217	256/405 (62%), Gaps = 2) nucleoside transporter Length = 406
Identities = 101/278 (36%), Positives = 160/278 (57%), Gaps = 1/278 (0%) splQ54087 LRPR_STREQ LEUCINE RICH PROTEIN pir  S39972 leucine-rich protein - Streptococcus equisimilis emb CAA51350.1  (X72832) leucine rich protein [Streptococcus equisimilis] prf  2009358B Leu-rich protein [Streptococcus equisimilis] Length = 282	Identities = 320/377 (84%), Positives = 359/377 (94%) splQ00752IMSMK_STRMU MULTIPLE SUGAR-BINDING TRANSPORT ATP-BINDING PROTEIN MSMK pir [E42400 ABC type transport system ATP-binding protein msmK [validated] - Streptococcus mutans gb AAA26938.1  (M77351) ATP-binding protein [Streptococcus mutans]	Identities = 342/399 (85%), Positives = 378/399 (94%) emb CAC21180.1  (AJ251799) putative phosphopentomutase [Streptococcus thermophilus]	Identities = 99/199 (49%), Positives = 140/199 (69%), Gaps = 1/199 (0%) splP43048 DEOC_MYCHO DEOXYRIBOSE-PHOSPHATE ALDOLASE (PHOSPHODEOXYRIBOALDOLASE) (DEOXYRIBOALDOLASE) pir  S42197 deoxyribose-phosphate aldolase (EC 4.1.2.4) - Mycoplasma hominis (SGC3) pir  S72522 deoxyribose-phosphate aldolase (EC 4.1.2.4) - Mycoplasma hominis (SGC3) deoxyribose aldolase (Mycoplasma hominis) (SGC3) emb CAA81646.1  (Z27121) deoxyribose aldolase [Mycoplasma hominis] Length = 217	Identities = 160/405 (39%), Positives = 256/405 (62%), Gaps = 8/405 (1%) dbj BAB05165.1  (AP001512) nucleoside transporter [Bacillus halodurans] Length = 406
47	06	91	. 25	62
Contig133 (9119-9970 m)	Contig133 (7885-9018 m)	Contig82 (6068-7264 p)	Contig82 (5330-6001 p)	Contig82 (4098-5300 p)
SA-1915.1	SA-1916.2	SA-1918.2	SA-1919.1	SA-1921.1
SeqID 987	SeqID 988	SeqID 989	SeqID 990	SeqID 991

ŏ	Contig82 (3298-4077 p) 63		Identities = 145/246 (58%), Positives = 171/246 (68%) splO83990 UDP_TREPA URIDINE PHOSPHORYLASE (UDRPASE) pir  F71251 probable uridine phosphorylase (udp) syphilis spirochete gb AAC65977.1  (AE001270) uridine phosphorylase (udp) [Treponema pallidum] Length = 258
Contig82 (24	Contig82 (2403-3140 m)	•	Identities = 50/171 (29%), Positives = 86/171 (50%), Gaps = 17/171 (9%) dbj BAB06113.1  (AP001515) transcriptional regulator (GntR family) [Bacillus halodurans] Length = 242
Contig82 (2094-2402 m)	94-2402 m) No Hits found	found	
Contig82 (803	803-1717 m) 45		Identities = 84/314 (26%), Positives = 140/314 (43%), Gaps = 28/314 (8%) pir  S55315 mucin (clone PGM-2A) - pig pir  I47141 gastric mucin (clone PGM-2A) - pig (fragment) gb AAC48526.1  (U10281) gastric mucin [Sus scrofa]
Contig82 (372-	(372-1994 p) 94	-	Identities = 471/539 (87%), Positives = 512/539 (94%), Gaps = 1/539 (0%) gb AAD23455.1  (AF117741) chaperonin GroEL [Streptococcus pneumoniae]
Contig82 (3-344 m)	44 m) No Hits found	found	
Contig82 (16-276 p)	-276 p) 81		Identities = 52/91 (57%), Positives = 75/91 (82%), Gaps = 1/91 (1%) gb AAD23454.1  (AF117741) cochaperonin GroES [Streptococcus pneumoniae]
Contig84 (8598	3598-9938 m) 63	m	Identities = 191/454 (42%), Positives = 289/454 (63%), Gaps = 17/454 (3%) dbj[BAB04579.1] (AP001510) BH0860~unknown conserved protein [Bacillus halodurans] Length = 458



SeqID 1000	SA-193.2	Contig136 (63475-63975 m)	64	Identities = 96/164 (58%), Positives = 125/164 (75%), Gaps = 1/164 (0%) sp[P42923]RL10_BACSU 50S RIBOSOMAL PROTEIN L10 (BL5) (COLD ACCLIMATIZATION PROTEIN) (CAP) (VEGETATIVE PROTEIN 300) (VEG300) pir[]D69695 ribosomal protein L10 (BL5) rplJ - Bacillus subtilis dbj BAA08840.1  (D50303) Ribosomal Protein L10 (Bacillus subtilis] emb CAB11880.1  (Z99104) ribosomal protein L10 (BL5) [Bacillus subtilis]
SeqID 1001	SA-1930.1	Contig84 (7772-8539 p)	44	Identities = 72/213 (33%), Positives = 115/213 (53%), Gaps = 11/213 (5%) pir  C70361 conserved hypothetical protein aq_701 - Aquifex aeolicus gb AAC06891.1  (AE000703) hypothetical protein [Aquifex aeolicus]
SeqID 1002	SA-1931.1	Contig84 (5832-7613 m)	73	Identities = 344/578 (59%), Positives = 446/578 (76%) splP14951 UVRC_BACSU EXCINUCLEASE ABC SUBUNIT C pir  A37192 excinuclease ABC; chain C - Bacillus subtilis gb AAA87316.1  (J03294) deoxyribodipyrimidine photolyase [Bacillus subtilis] emb CAA99578.1  (Z75208) deoxyribodipyrimidine photolyase [Bacillus subtilis] emb CAB14809.1  (Z99118) excinuclease ABC (subunit C) [Bacillus subtilis] Length = 598
SeqID 1003	SA-1932.2	Contig84 (3955-5790 p)	45	Identities = 151/585 (25%), Positives = 281/585 (47%), Gaps = 31/585 (5%) gb AAK06095.1 AE006429_13 (AE006429) UNKNOWN PROTEIN [Lactococcus lactis subsp. lactis] Length = 617
SeqID 1004	SA-1933.2	Contig84 (3237-3839 m)	65	Identities = 94/203 (46%), Positives = 134/203 (65%), Gaps = 3/203 (1%) gb AAK04278.1 AE006255_8 (AE006255)
SeqID 1005	SA-1934.2	Contig84 (1784-3190 m)	77	Identities = 311/474 (65%), Positives = 369/474 (77%), Gaps = 11/474 (2%) gb[AAC45369.1  (U78036) dipeptidase [Lactococcus lactis] Length = 472

	No Hits found	Contig103 (17660-17875 p)	SA-1941.1	SeqID 1011
Identities = 112/141 (79%), Positives = 124/141 (87%) splQ9ZA56 HPRK_STRMU HPR(SER) KINASE/PHOSPHATASE pir  T11568 probable HPr kinase (EC 2.7.1) - Streptococcus mutans gb AAC80172.1  (U75480) putative HPr(ser) kinase [Streptococcus mutans]	85	Contig103 (17991-18413 p)	SA-1940.1	SeqiD 1010
Identities = 83/123 (67%), Positives = 96/123 (77%), Gaps = 2/123 (1%) splP02394 RL7_BACSU 50S RIBOSOMAL PROTEIN L7/L12 (BL9) (A TYPE) (VEGETATIVE PROTEIN 341) (VEG341) pir  R5BS9 ribosomal protein L7/L12 - Bacillus subtilis emb CAB11881.1  (299104) ribosomal protein L12 (BL9) [Bacillus subtilis]	73	Contig136 (63046-63411 m)	SA-194.1	SeqID 1009
Identities = 27/63 (42%), Positives = 36/63 (56%) pir  T31110 extrace  ular matrix binding protein - Abiotrophia defectiva (fragment) gb AAD03320.1  (AF067776) extrace  ular matrix binding protein [Abiotrophia defectiva] Length = 2055	40	Contig84 (3-185 m)	SA-1939.1	SeqID 1008
Identities = 70/244 (28%), Positives = 103/244 (41%), Gaps = 31/244 (12%) pir[E72330 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD35886.1 AE001748_2 (AE001748) conserved hypothetical protein [Thermotoga maritima] Length = 233	39	Contig84 (354-1088 m)	SA-1937.1	SeqID 1007
Identities = 125/192 (65%), Positives = 163/192 (84%), Gaps = 1/192 (0%) splO86222 Y22A_HAEIN HYPOTHETICAL PROTEIN HI0220.2 gblAAC21888.1 (U32707) H. influenzae predicted coding region HI0220.2 [Haemophilus influenzae Rd] Length = 214	92	Contig84 (1103-1687 m)	SA-1935.1	SeqID 1006



SeqID 1012	SA-1942.1	Contig103 (17500-17763 p)	38	Identities = 21/63 (33%), Positives = 36/63 (56%), Gaps = 2/63 (3%) pir  E70043 hypothetical protein yvlC - Bacillus subtilis gb AAC67275.1  (AF017113) YvlC [Bacillus subtilis] emb CAB15516.1  (Z99121) yvlC [Bacillus subtilis] emb CAB15528.1  (Z99122) yvlC [Bacillus subtilis] Length = 65
SeqID 1013	SA-1943.1	Contig103 (16961-17419 p)	22	Identities = 63/146 (43%), Positives = 92/146 (62%), Gaps = 5/146 (3%) dbj BAA90855.1  (AB031213) YdcK [Bacillus halodurans] dbj BAB04251.1  (AP001508) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 151
SeqID 1014	SA-1944.1	Contig103 (14836-16998 p)		Identities = 379/729 (51%), Positives = 515/729 (69%), Gaps = 25/729 (3%) dbj BAB04250.1  (AP001508) unknown conserved protein [Bacillus halodurans] Length = 728
SeqID 1015	SA-1945.1	Contig103 (13688-14731 p)		Identities = 164/300 (54%), Positives = 217/300 (71%), Gaps = 1/300 (0%) gblAAB94650.1  (U96107) N5,N10-methylenetetrahydromethanopterin reductase homolog [Staphylococcus carnosus] Length = 301
SeqID 1016	SA-1946.1	Contig103 (13427-13555 m)	No Hits found	
SeqID 1017	SA-1949.1	Contig103 (12519-13334 m)	. 64	Identities = 141/283 (49%), Positives = 198/283 (69%), Gaps = 3/283 (1%) pir  E69759 hypothetical protein ycgR - Bacillus subtilis dbj  BAA08959.1  (D50453) homologue of unidentified protein of E. coli [Bacillus subtilis] emb  CAB12119.1  (299105) ycgR [Bacillus subtilis]
SeqID 1018	SA-195.1	Contig136 (62894-63439 p)	45	Identities = 38/157 (24%), Positives = 82/157 (52%), Gaps = 6/157 (3%) pir[ T39903 serine-rich protein - fission yeast (Schizosaccharomyces pombe) emb CAA22127.1  (AL033534) hypothetical serine-rich secreted protein [Schizosaccharomyces pombe] Length = 534
SeqID 1019	SA-1950.1	Contig103 (12179-12337 p)	No Hits found	

	ontig111 (16788-17714 m)	SA-1955.1 Contig111 (16788-17714 m)
778 m)	.contig111 (16137-16778 m)	SA-1956.1 Contig111 (16137-16



		<del>                                      </del>	$\overline{}$	Τ-	C		1
Identities = 85/335 (25%), Positives = 171/335 (50%), Gaps = 15/335 (4%) pir[ H72265 hypothetical protein TM1336 - Thermotoga maritima (strain MSB8) gb AAD36408 1 AE001788_3 (AE001788) permease, putative [Thermotoga maritima]	Identities = 75/279 (26%). Positives = 144/279 (50%) splP49330JRGG_STRGC RGG PROTEIN pirijA41898 positive regulator of glucosyltransferase and Spp phenotype rgg - Streptococcus gordonii gbjAAA26968.1j (M89776) rgg [Streptococcus gordonii]				Identities = 32/126 (25%), Positives = 52/126 (40%), Gaps = 20/126 (15%) pir]F71614 chromatinic RING finger DRING protein homolog PFB0440c - malaria parasite (Plasmodium falciparum) gbjAAC71877.1 (AE001395) chromatinic RING finger protein, DRING ortholog [Plasmodium falciparum]	Identities = 33/113 (29%), Positives = 53/113 (46%), Gaps = 19/113 (16%) pir  E71619 RAD2 endonuclease PFB0265c - malaria parasite (Plasmodium falciparum) gb AAC71842.1  (AE001383) RAD2 endonuclease [Plasmodium falciparum]	Identities = 155/698 (22%), Positives = 293/698 (41%), Gaps = 157/698 (22%) pir  F70175 rep helicase, single-stranded DNA-dependent ATPase (rep) homolog - Lyme disease spirochete gb AAC66967.1  (AE001162) rep helicase, single-stranded DNA-dependent ATPase (rep) [Borrelia burgdorferi] Length = 659
20	49	No Hits found	No Hits found	No Hits found	24	. 26	43
Contig111 (14894-15880 p)	Contig111 (13949-14821 p)	1 1	Contig136 (62632-62808 m)	Contig111 (12013-12432 p)	Contig111 (11221-11808 p)	Contig111 (10793-11221 p)	Contig136 (26286-28304 m)
SA-1957.1	SA-1958.1	SA-1959.1	SA-1960 1	SA-1961.1	SA-1962.2	SA-1963.2	SA-1966.1
SeqID 1026	SeqID 1027	SeqID 1028	SeaID 1030	SeqID 1031	SeqID 1032	SeqID 1033	SeqID 1034

SeqID 1035	SA-1967.1	Contig136 (25592-25951 m)	. SS	Identities = 61/118 (51%), Positives = 79/118 (66%), Gaps = 1/118 (0%) gb AAC98436.1  (L29324) unknown [Streptococcus pneumoniee] Length = 118
SeqID 1036	SA-1968.1	Contig136 (25217-25582 m)	92	Identities = 72/121 (59%), Positives = 95/121 (78%) gb AAC98435.1  (L29324) unknown [Streptococcus pneumoniae] Length = 121
SeqID 1037	SA-1969.1	Contig136 (23353-25230 m)	49	Identities = 241/416 (57%), Positives = 312/416 (74%), Gaps = 4/416 (0%) gb[AAC98434.1] (L29324) relaxase [Streptococcus pneumoniae]
SeqID 1038	SA-197.1	Contig136 (61820-62635 m)	8	Identities = 195/268 (72%), Positives = 233/268 (86%), Gaps = 1/268 (0%) gblAAD41248.1 AF106927_1 (AF106927) unknown [Streptococcus suis]
SeqID 1039	SA-1971.2	Contig136 (22443-23198 m)	77	Identities = 146/250 (58%), Positives = 197/250 (78%) splP26422 LACR_STRMU LACTOSE PHOSPHOTRANSFERASE SYSTEM REPRESSOR pir] B43258 regulatory protein lacR - Streptococcus mutans gb AAAZ6903.1  (M80797) lactose repressor [Streptococcus mutans] Length = 251
SeqID 1040	SA-1973.3	Contig117 (6819-7409 m)	54	Identities = 78/193 (40%), Positives = 111/193 (57%), Gaps = 14/193 (7%) gbjAAK04415.1 AE006268_11 (AE006268) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 191
SeqID 1041	SA-1974.2	Contig117 (5976-6734 p)	No Hits found	
SeqID 1042	SA-1975.2	Contig117 (5755-5973 p)	99	Identities = 28/60 (46%), Positives = 45/60 (74%) pir[IG75183 probable repressor protein PAB7155 - Pyrococcus abyssi (strain Orsay) emb[CAB49550.1] (AJ248284) repressor protein, putative [Pyrococcus abyssi]
SeqID 1043	SA-1976.2	Contig117 (5390-5713 p)	No Hits found	



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Identities = 70/258 (27%), Positives = 122/258 (47%), Gaps = 14/258 (5%) splP27129JRFAJ_ECOLI LIPOPOLYSACCHARIDE 1,2-GLUCOSYLTRANSFERASE pirl S47847 lipopolysaccharide glucosyltransferase I (EC 2.4.1.58) - Escherichia coli gb AAB18603.1  (U00039) UDP-D-Glucose:(galactosyl)lipopolysaccharide glucosyltransferase [Escherichia coli] gb AAC76650.1  (AE000440) UDP-D-Glucose:(galactosyl)lipopolysaccharide glucosyltransferase [Escherichia coli K12] Length = 338	Identities = 84/242 (34%), Positives = 147/242 (60%), Gaps = 16/242 (6%) dbj BAB07774.1  (AP001520) unknown conserved protein [Bacillus halodurans] Length = 236	Identities = 90/141 (63%), Positives = 102/141 (71%) gb AAC98422.1  (L29323) methyl transferase [Streptococcus pneumoniae] Length = 172	Identities = 482/885 (54%), Positives = 618/885 (69%), Gaps = 27/885 (3%) dbjjBAB04986.1  (AP001511) alanyl-tRNA synthetase [Bacillus halodurans] Length = 879	Identities = 43/144 (29%), Positives = 75/144 (51%), Gaps = 2/144 (1%) pir  G70079 hypothetical protein yxjl - Bacillus subtilis dbj BAA11710.1  (D83026) hypothetical [Bacillus subtilis] emb CAB15920.1  (299123) yxjl [Bacillus subtilis] Length = 162	Identities = 60/175 (34%), Positives = 103/175 (58%), Gaps = 6/175 (3%) gb AAK05744.1 AE006395_4 (AE006395) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 186
	,				
42	99	11	70	42	55
Contig117 (4126-4944 p)	Contig117 (3322-4038 p)	Contig136 (61161-61589 m)	Contig117 (622-3240 p)	Contig117 (121-606 p)	Contig128 (12351-12893 m)
SA-1977.1	SA-1978.1	SA-198.1	SA-1982.1	SA-1983.1	SA-1985.2
SeqiD 1044	SeqID 1045	SeqID 1046	SeqID 1047	SeqID 1048	SeqID 1049

SeqiD 1050	SA-1986.1	Contig 128 (9927-12275 m)	80	Identities = 321/790 (40%). Positives = 466/790 (58%), Gaps = 18/790 (2%) spjP94545jMUS2_BACSU MUTS2 PROTEIN pirjID69985 DNA mismatch repair protein homolog yshD - Bacillus subtilis emb[CAA99569.1] (Z75208) hypothetical protein [Bacillus subtilis] emb[CAB14818.1] (Z99118) similar to DNA mismatch repair protein [Bacillus subtilis]
SeqID 1051	SA-1987.1	Contig128 (9423-9770 m)	42	Identities = 42/143 (29%), Positives = 71/143 (49%), Gaps = 7/143 (4%) splQ57819 Y374_METJA HYPOTHETICAL PROTEIN MJ0374 pir  F64346 hypothetical protein MJ0374 - Methanococcus jannaschii gb AAB98363.1  (U67490) lipoprotein B (lppB) [Methanococcus jannaschii] Length = 330
SeqID 1052	SA-1988.1	Contig128 (9028-9360 m)	09	Identities = 64/100 (64%), Positives = 78/100 (78%), Gaps = 1/100 (1%) emb CAB40815.2  (AJ133006) thloredoxin [Listeria monocytogenes] Length = 103
SeqID 1053	SA-1989.1	Contig128 (8389-8982 p)	25	Identities = 27/67 (40%), Positives = 51/67 (75%) pirj  40868 hypothetical protein 3 nanH region - Clostridium perfringens emb CAA60798.1  (X87369) ORF3 [Clostridium perfringens] Length = 265
SeqID´1054	SA-199.1	Contig136 (60382-61734 m)	85	Identities = 322/448 (71%), Positives = 386/448 (85%), Gaps = 3/448 (0%) gb AAC98421.1  (L29323) methyl transferase
SeqID 1055	SA-1990.2	Contig128 (7088-8212 p)	. 28	Identities = 169/388 (43%), Positives = 224/388 (57%), Gaps = 26/388 (6%) gb AAK04939.1 AE006318_2 (AE006318) A/G-specific adenine glycosylase (EC 3.2.2.1) [Lactococcus lactis subsp. lactis] Length = 387
SeqID 1056	SA-1992.1	Contig128 (47161-47817 m)	19	Identities = 31/105 (29%), Positives = 51/105 (48%), Gaps = 4/105 (3%) emb CAB95931.1  (AL359989) galactose-1-phosphate uridylyltransferase [Streptomyces coelicolor A3(2)]

Contig128 (45990-47117 p) 65 27/410 (6%) db  BAB0 protein [Bacillu      Identities = 186/410 (4     Contig128 (45274-45810 m)	29 44 65 18 18 18 18 18 18 18 18 18 18 18 18 18 1	Identities = 186/410 (45%), Positives = 259/410 (62%), Gaps = 27/410 (6%) dbj BAB06998.1  (AP001518) unknown conserved protein [Bacillus halodurans]	Identities =_51/173_(29%), Positives = 87/173 (49%), Gaps = 20/173 (11%) emb CAA73267_1  (Y12736) orfX [Lactococcus lactis subsp. cremoris]	Identities = 29/87 (33%), Positives = 51/87 (58%), Gaps = 1/87 (1%) dbj BAB06137.1  (AP001515) DNA polymerase III (alpha subunit) [Bacillus halodurans]	Identities = 46/141 (32%), Positives = 71/141 (49%), Gaps = 11/141 (7%) splP32184 TIPA_STRCO TRANSCRIPTIONAL ACTIVATOR TIPA pir  S35354 tipA protein - Streptomyces lividans pir  T36339 transcription regulator - Streptomyces coelicolor gb AAB27737.1  (S64314) TipAL-AS=thiostrepton-pecific recognition protein(TipAL=transcriptional activator, ipAS=transcriptional activation modulator) [Streptomyces lividans, Peptide, 253 aa] emb CAB42766.1  (AL049841) transcriptional regulator [Streptomyces coelicolor A3(2)] Length = 253	Identities = 166/397 (41%), Positives = 255/397 (63%), Gaps =
	Contig128 (45990-47117 p)  Contig128 (44643-45251 m)  Contig128 (43791-44516 m)	27/410 (6%) dbj BAB06998.1  (AP00 protein [Bacillus halodurans]	Identities = 51/173 (29%), Posit 20/173 (11%) emb CAA73267.1 lactis subsp. cremoris]	Identities = 29/87 (33%), Positives	Identities = 46/141 (32%), Positives = 71/11/41 (7%) splP32184 TIPA_STRCO TIPA DITINA_STRCO TIPA DITINA_STRCO TIPA DITINA_STRCO TIPA DITINA DI	Identities = 166/397 (41%), Positives = 255/397 (63%), Gaps = 14/397 (3%) pir  H64571 cyclopropane-fatty-acyl-phospholipid
Contig128 (45990-47117 p)  Contig128 (44643-45810 m)  Contig128 (44643-45251 m)		65	44	24	87	
	SA-1993.1 SA-1996.1	Contig128-(45990-47117 p)	Contig128 (45274-45810 m)	Contig128 (44643-45251 m)	Contig128 (43791-44516 m)	

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Identities = 44/97 (45%), Positives = 60/97 (61%) sp P24281 YAAK_BACSU HYPOTHETICAL 11.8 KD PROTEIN	hypothetical protein yaaK - Bacillus subtilis emb CAA34878.1  (X17014) ORF107 [Bacillus subtilis] - dbj BAA05256.1  (D26185) unknown [Bacillus subtilis] emb CAB11796.1  (Z99104) similar to hypothetical proteins [Bacillus subtilis]  Length = 107			Identities = 37/161 (22%), Positives = 75/161 (45%), Gaps = 8/161 (4%) pir  S45089 hypothetical protein eta - Streptococcus pyogenes plasmid pDB101 pir  S68123 hypothetical protein eta - Streptococcus pyogenes plasmid pMD101 emb CAA47095.1  (X66468) orf eta [Streptococcus pyogenes] Length = 231	Identities = 69/143 (48%), Positives = 97/143 (67%), Gaps = 3/143 (2%) gb AAC98423.1 (L29323) unknown [Streptococcus pneumoniae]		Identities = 640/941 (68%), Positives = 784/941 (83%), Gaps = 3/941 (0%) splO34863 UVRA_BACSU EXCINUCLEASE ABC SUBUNIT A pir  F69729.excinuclease ABC chain A - Bacillus subtilis gb AAC67271.1  (AF017113) excinuclease ABC subunit A [Bacillus subtilis] emb CAB15533.1  (299122) excinuclease ABC (subunit A) [Bacillus subtilis]		Identities = 108/295 (36%), Positives = 174/295 (58%), Gaps = 4/295 (1%) pir  A75272 probable transport protein - Deinococcus radiodurans (strain R1) gb AAF12002.1 AE002075_6 (AE002075) transport protein, putative [Deinococcus radiodurans] Length = 312
:	44	No Hits found	No Hits found	50	59	No Hits found		No Hits found	90
	Contig128 (42071-42370 p)	Contig128 (41421-41759 m)	99 9 1		Contig128 (40742-41188 m)	Contig128 (151-2997 m)	Contig128 (3111-3782 m)	Contig128 (3807-4751 m)	
	SA-1998.1	SA-1999.1	SA-2.1	SA-20.1	SA-200.1	SA-2000.1	SA-2004.1	SA-2005.1	SA-2006.1
:	SeqID 1062	SeqID 1063	SeqID 1064	SeqID 1065	SeqID 1066	SeqID 1067	SeqID 1068	SeqID 1069	SeqID 1070



Identities = 52/79 (65%), Positives = 64/79 (80%) pir  S66013 ribosomal protein S18 (rpsR) - Bacillus subtilis dbj BAA05219.1  (D26185) ribosomal protein S18 [Bacillus subtilis] emb CAB16126.1  (Z99124) ribosomal protein S18 [Bacillus subtilis] Length = 81	Identities = 136/163 (83%), Positives = 149/163 (90%)   gb AAF98351.1  (AF280767) single strand binding protein   Streptococcus pyogenes]	Identities = 72/96 (75%), Positives = 81/96 (84%), Gaps = 1/96 (1%) gb AAK06289.1 AE006448_7 (AE006448) 30S ribosomal protein S6 [Lactococcus lactis subsp. lactis] Length = 97	Identities = 52/93 (55%), Positives = 68/93 (72%)   gb AAC14608.1  (U95840) transmembrane protein Tmp5   [Lactococcus lactis]		Identities = 92/204 (45%), Positives = 101/204 (49%), Gaps = 1/204 (0%) pir  H72504 hypothetical protein APE2014 - Aeropyrum pernix (strain K1) dbj BAA81024.1  (AP000063) 280aa long hypothetical protein [Aeropyrum pernix] Length = 280		Identities = 490/652 (75%), Positives = 561/652 (85%), Gaps = 5/652 (0%) gb AAC16243.2  (AF061748) cell division protein FtsH   Streptococcus pneumoniae] Length = 652
73	06	. 42	64	No Hits found	40	No Hits found	82
Contig128 (4921-5160 m)	Contig128 (5205-5696 m)	Contig128 (5708-5995 m)	Contig113 (25825-26142 p)	Contig113 (25470-25775 m)	Contig113 (22823-23569 m)	Contig113 (22538-22783 m)	Contig113 (22413-24389 p)
SA-2007.1	SA-2008.2	SA-2009.2	SA-2010.1	SA-2012.1	SA-2013.1	SA-2014.1	SA-2015.2
SeqID 1071	SeqID 1072	SeqID 1073	SeqID 1074	SeqID 1075	SeqID 1076	SeqID 1077	SeqID 1078

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Identities = 121/179 (67%), Positives = 152/179 (84%), Gaps = 1/179 (0%) splQ02522 HPRT_LACLA HYPOXANTHINE-GUANINE PHOSPHORIBOSYLTRANSFERASE (HGPRT) (HGPRTASE) pir  S30100 hypoxanthine phosphoribosyltransferase (EC 2.4.2.8) - Lactococcus lactis emb CAA48876.1  (X69123) hypoxanthine guanine phosphoribosyltransferase (EC 2.4.2.8) - Lactococcus lactis emb CAA47404.1  (X67015) hypoxanthine phosphoribosyltransferase (EC 2.4.2.8) [Lactococcus lactis] gb AAK04118.1 AE006241_7 (AE006241) hypoxantine-guanine phosphoribosyltransferase (EC 2.4.2.8) [Lactococcus lactis subsp. lactis] prf  1905381A hypoxanthine guanine phosphoribosyltransferase [Lactococcus lactis] Length = 183	Identities = 142/418 (33%), Positives = 230/418 (54%), Gaps = 21/418 (5%) gb AAK04117.1 AE006241_6 (AE006241) cell cycle protein MesJ [Lactococcus lactis subsp. lactis]  Length = 423	Identities = 77/284 (27%), Positives = 117/284 (41%), Gaps = 54/284 (19%) prf[ 1714238A beta lactamase mutant S-3P [Staphylococcus aureus]		Identities = 159/334 (47%), Positives = 236/334 (70%), Gaps = 15/334 (4%) gb AAK05774.1 AE006398_5 (AE006398) transcriptional regulator [Lactococcus lactis subsp. lactis]	Identities = 121/418 (28%), Positives = 190/418 (44%), Gaps = 43/418 (10%) splP29850 MALX_STRPN   MALTOSE/MALTODEXTRIN-BINDING PROTEIN PRECURSOR pir  S32569 malX protein - Streptococcus pneumoniae gb AAA26925.1  (L08611) MalX [Streptococcus pneumoniae]   Length = 423
æ	53	56	No Hits found	- 88 90	42
Contig113 (21848-22390 p)	Contig113 (20569-21843 p)	Contig113 (19401-20567 p)	Contig136 (59573-59959 m)	Contig93 (6702-7730 p)	Contig93 (5199-6446 m)
SA-2017.2	SA-2018.2	SA-2019.2	SA-202.1	SA-2022.2	SA-2024.1
SeqID 1079	SeqID 1080	SeqID 1081	SeqID 1082	SeqID 1083	SeqiD 1084



Identities = 174/428 (40%), Positives = 263/428 (60%), Gaps = 24.2025.1   Contig93 (3731-5071 m)   56   0xytoca emb[CAA60005.1  (X86014) cymF [Klebsiella oxytoca]   Length = 427	Identities = 109/269 (40%), Positives = 171/269 (63%), Gaps = 5/269 (1%) pir  S63616 malG protein homolog cymG - Klebsiella oxytoca emb CAA60006.1  (X86014) cymG [Klebsiella oxytoca]	Identities = 259/485 (53%), Positives = 353/485 (72%), Gaps = 9/485 (1%) sp P94408 YCLF_BACSU HYPOTHETICAL 53.3 KD PROTEIN IN SFP-GERKA INTERGENIC REGION pir] C69762 di-tripeptide ABC transporter (membrane pr) homolog yclF - Bacillus subtilis dbj BAA09000.1  (D50453) homologue of Di-tripeptide transporter Dtp of L. lactis [Bacillus subtilis] emb CAB12175.1  (299106) similar to di-tripeptide ABC transporter (membrane protein) [Bacillus subtilis] transporter (membrane protein) [Bacillus subtilis]	Identities = 37/114 (32%), Positives = 66/114 (57%), Gaps = 10/114 (8%) dbj BAB07289.1  (AP001519) mutator MutT protein [Bacillus halodurans] Length = 159	SA-203.1 Contig136 (5	SA-2030.1	SA-2031.1 Contig93	Identities = 73/188 (38%), Positives = 108/188 (56%), Gaps = 108/2033.2   Contig123 (38769-39380 p)   52   11/188 (5%) dbj BAA21095.1   (D88438) repB [Lactobacillus acidophilus]   Length = 193	SA-2034.1 Contig123 (37952-38569 p) No Hits found	SA-2035.2
SA-2025.1	SA-2026.	SA-2028.	SA-2029.	SA-203.1	SA-2030.	SA-2031.	SA-2033.2	SA-2034.1	SA-2035.2
SeqiD 1085	SeqID 1086	SeqID 1087	SeqID 1088	SeqID 1089	SeqID 1090	SeqID 1091	SeqID 1092	SeqID 1093	SeaID 1094

SeqID 1095 SeqID 1096	SA-2037.2 SA-2038.1	Contig123 (36547-36867 p) Contig123 (35896-36345 p)	No Hits found No Hits found	
SeqID 1097	SA-2040.1	Contig 123 (34160-35575 p)	No Hits found	
SeqID 1098	SA-2041.2	Contig123 (33771-34109 p)	09	Identities = 50/110 (45%), Positives = 76/110 (68%) ref[NP_054018.1  CadX (Staphylococcus lugdunensis] gb AAB18271.1  (U74623) CadX (Staphylococcus lugdunensis] Length = 115
SeqID 1099	SA-2042.2	Contig123 (33133-33759 p)	26	Identities = 198/209 (94%), Positives = 203/209 (96%) pir  E81967 cadmium resistance protein NMA0496 [imported] - Neisseria meningitidis (group A strain Z2491) emb CAB83789.1  (AL162753) cadmium resistance protein [Neisseria meningitidis Z2491] Length = 213
SeqID 1100	SA-2045.2	Contig132 (41522-42709 p)	59	Identities = 171/393 (43%), Positives = 256/393 (64%), Gaps = 2/393 (0%) gb AAK06200.1 AE006438_14 (AE006438) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 400
SeqID 1101	SA-2047.1	Contig132 (39834-41324 p)	51	Identities = 186/500 (37%), Positives = 258/500 (51%), Gaps = 67/500 (13%) emb CAB95221.1  (AL359773) possible threonine synthase [Leishmania major]
SeqID 1102	SA-2048.1	Contig132 (38697-39713 p)	92	Identities = 238/340 (70%), Positives = 276/340 (81%), Gaps = 2/340 (0%) pir  H81186 alcohol dehydrogenase, propanolpreferring NMB0546 [imported] - Neisseria meningitidis (group B strain MD58) gb AAF40975.1  (AE002410) alcoholdehydrogenase, propanol-preferring [Neisseriameningitidis MC58] Length = 348
SeqID 1103	SA-205.1	Contig136 (58759-59349 m)	No Hits found	
SeqID 1104	SA-2050.2	Contig132 (35876-38518 p)	84	Identities = 658/873 (75%), Positives = 760/873 (86%), Gaps = 2/873 (0%) gb AAK03537.1  (AE006181) Adh2 [Pasteurella multocida] Length = 875
SeqID 1105	SA-2051.1	Contig129 (2-493 m)	28	Identities = 68/155 (43%), Positives = 98/155 (62%) gb[AAF13747.1 AF117351_4 (AF117351) unknown [Zymomonas mobilis] Length = 236

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			ļ	Identities = 227/341 (66%), Positives = 278/341 (80%)
SeqID 1106	SA-2052.1	Contig 129 (646-1674 p)	08	spi032034jqUEA_BACSU S-ADENOSYLME I HIONINE: I RNA RIBOSYLTRANSFERASE-ISOMERASE (QUEUOSINE BIOSYNTHESIS PROTEIN QUEA) pirjlA69688 S-adenosylmethioninetRNA ribosylfransferase-isomerase (EC 5.4.99) queA [similarity] - Bacillus subtilis emb CAB14732.1  (Z99118) S-adenosylmethionine tRNA ribosyltransferase [Bacillus subtilis] emb CAB75332.1  (Y15896) QueA protein [Bacillus subtilis]
SeqID 1107	SA-2053.1	Contig129 (1761-2198 p)	54	Identities = 56/145 (38%), Positives = 86/145 (58%), Gaps = 2/145 (1%) emb CAA73494.1  (Y13052) ORF145 [Staphylococcus sciuri]
SeqID 1108 :	SA-2054.1	Contig129 (2253-3317 m)	No Hits found	
SeqID 1109	SA-2056.1	Contig129 (3418-4674 m)	74	Identities = 232/416 (55%), Positives = 314/416 (74%), Gaps = 3/416 (0%) splQ57493 Y092_HAEIN HYPOTHETICAL PROTEIN H10092 pir ID64142 hypothetical protein H10092 - Haemophilus influenzae (strain Rd KW20) gb AAC21770.1  (U32694) H. influenzae predicted coding region H10092 [Haemophilus influenzae Rd] Length = 419
SeqID 1110	SA-2059.2	Contig129 (4699-5841 m)	59	Identities = 177/367 (48%), Positives = 235/367 (63%), Gaps = 2/367 (0%) gb AAG58254.1 AE005541_6 (AE005541) orf, hypothetical protein [Escherichia coli O157:H7]  387
SeqID 1111	SA-206.1	Contig 136 (58190-58678 m)	No Hits found	
SeqID 1112	SA-2060.1	Contig129 (6008-7102 p)	90	Identities = 94/370 (25%), Positives = 184/370 (49%), Gaps = 13/370 (3%) dbj BAB06450.1  (AP001516) unknown conserved protein [Bacillus halodurans]
SeqID 1113	SA-2061.2	Contig129 (7171-8598 m)	80	Identities = 334/475 (70%), Positives = 392/475 (82%), Gaps = 8/475 (1%) gbJAAF89979.1 AF206272_5 (AF206272) betaglucosidase [Streptococcus mutans] Length = 479



SeqID 1114	SA-2062.2	Contig132 (47810-48070 p)	96	Identities = 82/86 (95%), Positives = 83/86 (96%) -sp[@9WW03]RS17_STRPN 30S RIBOSOMAL PROTEIN :S17— gb[AAD33265.1]AF126059_6 (AF126059) RpS17 [Streptococcus pneumoniae] gb[AAD33274.1] (AF126060) RpS17[Streptococcus pneumoniae] gb[AAD33283.1] (AF126061) RpS17 [Streptococcus pneumoniae]
SeqID 1115	SA-2063.2	Contig132 (47578-47784 p)	88	Identities = 58/68 (85%), Positives = 64/68 (93%) sp[Q9VVVV8]RL29_STRPN 50S RIBOSOMAL PROTEIN L29 gb]AAD33264.1 AF126059_5 (AF126059) RpL29 [Streptococcus pneumoniae] gb]AAD33273.1  (AF126060) RpL29 (Streptococcus pneumoniae] gb]AAD33282.1  (AF126061) RpL29 [Streptococcus pneumoniae]
SeqID 1116	SA-2065.1	Contig132 (47155-47568 p)	86	Identities = 135/137 (98%), Positives = 137/137 (99%) splQ9X5K1 RL16_STRPN 50S RIBOSOMAL PROTEIN L16 gb AAD33263.1 AF126059_4 (AF126059) RpL16 [Streptococcus pneumoniae] Length = 137
SeqID 1117	SA-2066.1	Contig132 (46498-47151 p)	91	Identities = 200/208 (96%), Positives = 203/208 (97%) sp[Q9WW37 RS3_STRPN 30S RIBOSOMAL PROTEIN S3 gb]AAD33262.1 AF126059_3 (AF126059) RpS3 [Streptococcus pneumoniae] gb AAD33271.1  (AF126060) RpS3 [Streptococcus pneumoniae] gb AAD33280.1  (AF126061) RpS3 [Streptococcus pneumoniae]
SeqID 1118	SA-2067.1	Contig132 (46141-46485 p)	68	Identities = 99/114 (86%), Positives = 106/114 (92%) splQ9WVU5[RL22_STRPN 50S RIBOSOMAL PROTEIN L22 gb AAD33261.1 AF126059_2 (AF126059) RpL22 [Streptococcus pneumoniae] gb AAD33270.1  (AF126060) RpL22 [Streptococcus pneumoniae] gb AAD33279.1  (AF126061) RpL22 [Streptococcus pneumoniae]

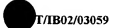
SeqID 1119	SA-2069.1	Contig132 (45847-46125 p)	98	Identities = 92/93 (98%). Positives = 93/93 (99%) sp[Q9WW12]RS19_STRPN 30S RIBOSOMAL PROTEIN S19 gb[AAD33260 1 AF126059_1 (AF126059) RpS19 (Streptococcus pneumoniae] gb[AAD33269 1  (AF126060) RpS19 (Streptococcus pneumoniae] gb[AAD33269 1 (AF126061) RpS19 (Streptococcus pneumoniae] gb[AAD33278.1  (AF126061) RpS19 (Streptococcus pneumoniae]
SeqID 1120	SA-207.1	Contig136 (56373-58190 m)	47	Identities = 183/492 (37%), Positives = 292/492 (59%), Gaps = 30/492 (6%) gblAAF72343.1 AF192329_4 (AF192329) TrsK-like protein [Enterococcus faecalis] Length = 564
SeqID 1121	SA-2071.1	Contig132 (44915-45748 p)	84	Identities = 207/277 (74%), Positives = 239/277 (85%) gb AAC45959.1  (U43929) L2 [Bacillus subtilis] Length = 277
SeqID 1122	SA-2072.1	Contig132 (44601-44897 p)	60	Identities = 56/92 (60%), Positives = 67/92 (71%), Gaps = 1/92 (1%) splQ9Z9L2 RL23_BACHD 50S RIBOSOMAL PROTEIN L23 pir  T44385 ribosomal protein L23 [imported] - Bacillus halodurans dbj BAA75273.1  (AB017508) rpIW homologue (identity of 71 to B. subtilis%) [Bacillus halodurans] dbj BAB03855.1  (AP001507) ribosomal protein L23 [Bacillus halodurans]
SeqID 1123	SA-2073.1	Contig132 (43978-44601 p)	73	Identities = 130/207 (62%), Positives = 160/207 (76%) splP42921 RL4_BACSU 50S RIBOSOMAL PROTEIN L4 pir  H69694 ribosomal protein L4 rpID - Bacillus subtilis dbj BAA08832.1  (D50302) Ribosomal Protein L4 [Bacillus subtilis] gb AAC45957.1  (U43929) L4 [Bacillus subtilis] emb CAB11893.1  (299104) ribosomal protein L4 [Bacillus subtilis]
SeqID 1124	SA-2074.2	Contig132 (43328-43954 p)	. 85	Identities = 157/208 (75%), Positives = 180/208 (86%), Gaps = 2/208 (0%) splP42920 RL3_BACSU 50S RIBOSOMAL PROTEIN L3 (BL3) pir  G69694 ribosomal protein L3 (BL3) rplC - Bacillus subtilis gb AAC45956.1  (U43929) L3 [Bacillus subtilis] emb CAB11892.1  (Z99104) ribosomal protein L3 (BL3) [Bacillus subtilis]



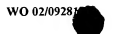
Identities = 98/102 (96%), Positives = 102/102 (99%)     sp P48853 RS10_STRMU 30S RIBOSOMAL PROTEIN S10     gb AAB46363.1  (L29637) S10 ribosomal protein [Streptococcus mutans]   Length = 102		Identities = 241/259 (93%), Positives = 248/259 (95%) emb CAB90834.1  (AJ250837) putative transposase [Streptococcus dysgalactiae] Length = 259	Identities = 93/96 (96%), Positives = 94/96 (97%) emb CAB90833.1  (AJ250837) hypothetical protein [Streptococcus dysgalactiae]			Identities = 1141/1150 (99%), Positives = 1142/1150 (99%)   gb AAB17762.1  (U56908) SCPB [Streptococcus agalactiae]   Length = 1150	Identities = 322/472 (68%), Positives = 378/472 (79%) pir  S68598 sucrose-6-phosphate hydrolase ScrB - Streptococcus sobrinus (strain 6715) Length = 479	Identities = 225/320 (70%), Positives = 273/320 (85%) splQ54430 SCRR_STRMU SUCROSE OPERON REPRESSOR (SCR OPERON REGULATORY PROTEIN) gb AAC31628.1  (U46902) ScrR [Streptococcus mutans] Length = 320	Identities = 51/129 (39%), Positives = 82/129 (63%), Gaps = 9/129 (6%) spIP54520 NUSB_BACSU N UTILIZATION SUBSTANCE PROTEIN B HOMOLOG (NUSB PROTEIN) pir[ F69960 transcription termination factor nusB homolog yqhZ [similarity] - Bacillus subtilis dbj BAA12571.1  (D84432) YqhZ [Bacillus subtilis] emb CAB14363.1  (Z99116) similar to transcription termination [Bacillus subtilis]
94	No Hits found	8	98	No Hits found	No Hits found	86	74	84	99
Contig132 (42915-43223 p)	Contig132 (42879-43280 m)	Contig92 (5603-6382 p)	Contig92 (5277-5567 p)	Contig92 (4012-5049 m)	Contig136 (56111-56353 m)	Contig92 (24-3476 m)	Contig102 (16409-17926 m)	Contig102 (15445-16407 m)	Contig102 (14924-15358 p)
SA-2075.2	SA-2076.2	SA-2077.1	SA-2078.1	SA-2079.1	SA-208.1	SA-2082.1	SA-2083.2	SA-2084.1	SA-2085.1
SeqID 1.125 -	SeqID 1126	SeqID 1127	SeqID 1128	SeqID 1129	SeqID 1130	SeqID 1131	SeqID 1132	SeqID 1133	SeqiD 1134

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SeqID 1135	SA-2086.1	- Contig102 (14542-14931 p)	53	Identities = 42/107 (39%), Positives = 70/107 (65%), Gaps = 4/107 (3%)-dbjlBAB06505.1  (AP001516) unknown conserved protein [Bacillus halodurans] Length = 132
SeqID 1136	SA-2087.1	Contig102 (13893-14453 p)	64	Identities = .89/186 (47%), Positives = 120/186 (63%), Gaps = 1/186 (0%) spIP49778 EFP_BACSU ELONGATION FACTOR P (EF-P) pir  A69620 translation elongation factor EF-P efp - Bacillus subtilis dbj BAA12558.1  (D84432) YqhU [Bacillus subtilis] emb CAB14376.1  (299116) elongation factor P [Bacillus subtilis]
SeqID 1137	SA-2090.1	Contig102 (10993-12714 p)	52	Identities = 174/568 (30%), Positives = 300/568 (52%), Gaps = 7/568 (1%) gb AAD10394.1  (U46488) NrpB [Proteus mirabilis] Length = 575
SeqID 1138	SA-2091.1	Contig 102 (10920-11162 m)	No Hits found	
SeqID 1139	SA-2092.2	Contig102 (9291-11003 p)	90	Identities = 186/583 (31%), Positives = 305/583 (51%), Gaps = 14/583 (2%) gb AAD10393.1  (U46488) NrpA [Proteus mirabilis] Length = 588
SeqID 1140	SA-2095.2	Contig119 (7199-8548 m)	06	Identities = 377/449 (83%), Positives = 414/449 (91%) pir  T51720 glucose-6-phosphate Isomerase (EC 5.3.1.9) [imported] - Streptococcus mutans gb AAD33517.1 AF132127_2 (AF132127) glucose-6-phosphate isomerase [Streptococcus mutans] Length = 449
SeqID 1141	SA-2096.1	Contig119 (6350-6877 m)	71	Identities = 96/173 (55%), Positives = 129/173 (74%) dbj BAA28715.1  (AB001562) hypothetical protein [Streptococcus mutans] Length = 178
SeqID 1142	SA-2097.1	Contig119 (5682-6359 m)	72	Identities = 126/218 (57%), Positives = 166/218 (75%) emb CAB90755.1  (AJ400707) hypothetical protein [Streptococcus uberis]



SeqID 1143	SA-2099.1	Contig119 (4507-5550 m)	09	Identities = 148/349 (42%), Posttives = 223/349 (63%), Gaps = 16/349 (4%) spiO05252jYUFN_BACSU HYPOTHETICAL LIPOPROTEIN YUFN PRECURSOR pri[C70009 ABC transporter (hoprotein) homolog yufN - Bacillus subtilis emb[CAB07936 1] (293937) unknown [Bacillus subtilis] emb[CAB15143 1] (299120) similar to ABC transporter (hoprotein) [Bacillus subtilis]
SeqID 1144	SA-21.1	Contig 137 (24502-24849 p)	No Hits found	
SeqID 1145	SA-210.1	Contig136 (55240-56094 m)	38	Identities = 56/262 (21%), Positives = 121/262 (45%), Gaps = 28/262 (10%) gb AAF72344.1 AF192329_5 (AF192329) unknown [Enterococcus faecalls] Length = 287
SeqID 1146	SA-2100.1	Contig119 (3517-4416 p)	63	Identities = 263/299 (87%), Positives = 287/299 (95%) dbj BAA28714.1  (AB001562) glucose-1-phosphate uridylyltransferase [Streptococcus mutans] 306
SeqiD 1147	SA-2101.1	Contig119 (2464-3480 p)	99	identities = 177/333 (53%), Positives = 241/333 (72%) sp P46919 GPDA_BACSU GLYCEROL-3-PHOSPHATE DEHYDROGENASE [NAD(P)+] (NAD(P)H-DEPENDENT DIHYDROXYACETONE-PHOSPHATE REDUCTASE) gb AAA86746.1  (U32164) NAD(P)H-dependent dihydroxyacetone-phosphate reductase [Bacillus subtilis] Length = 345
SeqID 1148	SA-2102.1	Contig119 (1965-2294 m)	45	Identities = 52/109 (47%), Positives = 77/109 (69%), Gaps = 1/109 (0%) pdb[1A6F] Rnase P Protein From Bacillus Subtitis Length = 119
SeqID 1149	SA-2103.2	Contig119 (1137-1952 m)	7.1	Identities = 149/267 (55%), Positives = 197/267 (72%), Gaps = 3/267 (1%) gb AAK04227.1 AE006251_3 (AE006251) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 269
SeqID 1150	SA-2104.2	Contig133 (64785-66146 m)	39	Identities = 100/451 (22%), Positives = 180/451 (39%), Gaps = 81/451 (17%) emb CAB39034.1  (AL034559) hypothetical protein, PFC0940c [Plasmodium falciparum] Length = 806
SeqID 1151	SA-2105.1	Contig133 (66147-67238 m)	No Hits found	



m) 50	No Hits found	D) No Hits found	m) No Hits found	p) No Hits found	Identities = 35/91 (38%), Positives = 51/91 (55%)   sp P34159 YHS1_CLOAB HYPOTHETICAL 11.0 KD PROTEIN   IN HSP18 3 REGION (ORFA1) emb CAA46375.1  (X65276)   ORFA1 (Clostridium acetobutylicum)   Length = 96	p) No Hits found	p) No Hits found	p) No Hits found	m) No Hits found	m) No Hits found	Identities = 260/311 (83%), Positives = 283/311 (90%) sp P95765 PPAC_STRGC PROBABLE MANGANESE- DEPENDENT INORGANIC PYROPHOSPHATASE (PYROPHOSPHATE PHOSPHO-HYDROLASE) (PPASE) gb AAB39104.1  (U57759) intrageneric coaggregation-relevant adhesin [Streptococcus gordonii] Length = 311	Identities = 261/793 (32%), Positives = 436/793 (54%), Gaps = 36/793 (4%) gb AAF72347.1 AF192329_8 (AF192329) TrsE-like protein [Enterococcus faecalis] Length = 799	Identities = 185/260 (71%), Positives = 218/260 (83%)  splO68575 PFLA_STRMU PYRUVATE FORMATE-LYASE
Contig133 (67478-68422 m)	Contig133 (68486-68815 m)	Contig 133 (69774-70190 p)	Contig136 (54825-55058 m)	Contig133 (70239-70367 p)	Contig133 (70808-71098 p)	Contig133 (71260-71571 p)	Contig133 (71861-72589 p)	Contig 133 (72806-73078 p)	Contig133 (73197-73502 m)	Contig133 (73790-74143 m)	Contig118 (5080-6015 p)	Contig136 (52523-54868 m)	Contig118 (4175-4963 p)
- SA-2106.1	SA-2107.1	SA-2109.1	SA-211.1	SA-2110.1	SA-2112.1	SA-2113.1	SA-2115.1	SA-2116.1	SA-2117.1	SA-2118.2	SA-2119.2	SA-212.1	SA-2120.1
SeqID 1152 -	SeaID 1153	SeqID 1154	SeqID 1155	SeqID 1156	SeqID 1157	SeqID 1158	SeqID 1159	SeqID 1160	SeqID 1161	SeqID 1162	SeqID 1163	SeqID 1164	SeqiD 1165



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Identities = 349/445 (78%), Positives = 404/445 (90%), Gaps	Identities = 80/184 (43%), Positives = 119/184 (64%), Gaps = 3/184 (1%) pir] G81942 hypothetical protein NMA0960 [imported] Neisseria meningitidis (group A strain Z2491) emb CAB84230.1  (AL162754) hypothetical protein NMA0960 [Neisseria meningitidis] Length = 188	Identities = 161/302 (53%), Positives = 220/302 (72%), Gaps = 4/302 (1%) pir  D69999 conserved hypothetical protein ytqA - Bacillus subtilis gb AAC00380.1  (AF008220) YtqA [Bacillus subtilis] emb CAB15026.1  (Z99119) similar to hypothetical proteins [Bacillus subtilis] Length = 322	Identities = 62/159 (38%), Positives = 93/159 (57%), Gaps = 3/159 (1%) pir  S32217 hypothetical protein 2 - Bacillus megaterium emb CAA79986.1  (Z21972) ORF2 [Bacillus megaterium] Length = 216	Identities = 33/113 (29%), Positives = 62/113 (54%) sp P50726 YPAA_BACSU HYPOTHETICAL 20.5 KD PROTEIN IN SERA-FER INTERGENIC REGION pir  E69932 hypothetical protein ypaA - Bacillus subtilis gb AAC83944.1  (L47648) putative [Bacillus subtilis] emb CAB14237.1  (299116) ypaA [Bacillus subtilis]	Identities = 44/74 (59%), Positives = 53/74 (71%) pir  T44088 probable transposase [imported] - Staphylococcus aureus Length = 74
06	62	69	42	48	51
- Contig118-(2773-4107 p)	Contig118 (2027-2593 p)	Contig118 (1099-2034 p)	Contig118 (365-1006 p)	Contig1.18 (1-384 p)	Contig92 (6606-6896 p)
SA-2122.1	SA-2123.1	SA-2124.1	SA-2125.1	SA-2126.1	SA-2128.2
SeqID 1166	SeqID 1167	SeqID 1168	SeqID 1169	SeqID 1170	SeqID 1171



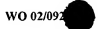
ss] cis ORFB 96	ocillus secis]	)11 cus	ictiae ctiae]	ctiae ae]	Gaps = VncR 06399) Length =
3%), Gap: ss anthraci lilus anthra U30714) C NFB [Bac 04) pXO11	3 (66%) pir[[H591] cis virulenc ORFB [Ba cillus anthr	b) pir  T09( pyogenes Streptococ	322 (98%) ccus agala ccus agala	306 (99%) ccus agala ıs agalact	(80%), Ga 40356) Vn II (AJ0063 Len
%), Positives = 86/135 (63%), Gaps = 52792.1  pXO1-96 [Bacillus anthracis] al protein pXO1-96 - Bacillus anthracis pXO1 gb AA74027.1  (U30714) ORFB AAA74029.1  (U30715) ORFB [Bacillus OO.1 AAD32400 (AF065404) pXO1-96 thracis] Length = 274	Identities = 45/98 (45%), Positives = 66/98 (66%) P_052792.1  pXO1-96 [Bacillus anthracis] pir  H59102 thetical protein pXO1-96 - Bacillus anthracis virulence pXO1 gb AAA74027.1  (U30714) ORFB [Bacillus is] gb AAA74029.1  (U30715) ORFB [Bacillus anthracis] AD32400.1 AAD32400 (AF065404) pXO1-96 [Bacillus anthracis] Length = 274	ss = 69/72 (95% Streptococcus 26542) TnpA [ Length = 364	/es = 816/6 Streptoco [Streptoco	ves = 304// Streptococitreptococitreptococci	= 176/218 56_1 (AF1 AB54582.1 eumoniae]
Positives = 12.1 pXO1 otein pXO 1 gb AAA 11 qb AAD3240 qcis]	5%), Positius (Bacillus 1-96 - Bacillus A74027.1  (U30715) (AF065,00)	ositives = or InpA - Stre	(98%), Positiv 2.4K protein - 33) unknown [ Length = 822	(99%), Positiv 5 [validated] - 5333	Positives: .1 AF1403 ae] emb C coccus pne
135 (47 INP_06 othetic is] gb  AD324 Illus an	= 45/98 (48 1.1 px01-6 otein px0 (O1 gb AA A74029.1 1 AAD324 anthracis]	72 (95%), Finsposase \text{AB92607.} pyogenes]	309/822 (9) hetical 92.4 AF062533	303/306 (9) otein Imb [ I (AF0625:	218 (60%), AAD25108 pneumoni or [Strepto
Identities = 64/135 (47%), Positives = 86/135 (63%), Gaps = 1/135 (0%) ref[NP_052792.1  pXO1-96 [Bacillus anthracis] pirl H59102 hypothetical protein pXO1-96 - Bacillus anthracis virulence plasmid pXO1 gb AAA74027.1  (U30714) ORFI [Bacillus anthracis] gb AAA74029.1  (U30715) ORFB [Bacillus anthracis] gb AAD32400.1 AAD32400 (AF065404) pXO1-96 [Bacillus anthracis]	dentitie 0527 thetical is] gb[/	Identities = 69/72 (95%), Positives = 69/72 (95%) pir  T09011 probable transposase TnpA - Streptococcus pyogenes (fragment) gb AAB92607.1  (AF026542) TnpA (Streptococcus pyogenes] Length = 364	Identities = 809/822 (98%), Positives = 816/822 (98%) pir  T46758 hypothetical 92.4K protein - Streptococcus agalactiae gb AAD13797.1  (AF062533) unknown [Streptococcus agalactiae] Length = 822	Identities = 303/306 (99%), Positives = 304/306 (99%) pir  T46757 lipoprotein Imb [validated] - Streptococcus agalactiae gb AAD13796.1  (AF062533) Lmb [Streptococcus agalactiae] Length = 306	Identities = 131/218 (60%), Positives = 176/218 (80%), Gaps: 1/218 (0%) gb AAD25108.1 AF140356_1 (AF140356) VncR [Streptococcus pneumoniae] emb CAB54582.1  (AJ006399) response regulator [Streptococcus pneumoniae] Length 218
Ident 1/13 pirl H virulenc (Bacill anthr	refINF hypol plasmid anthraci	Ident pr (fragn	lc pir  T46 gb AAD	lc pir  T46 gb A	Identi 1/21 [Stre respor
09	. 62	74	86	66	79
34 p)	38 b)	17 p)	84 m)	317 m)	3302 p)
(7096-7434 p)	(7421-7738 p)	(7784-8017 p)	Contig92 (8216-10684 m)	10697-11617 m)	(98649-99302 p)
Contig92	Contig92	Contig92	Contig92 (	Contig92 (1	Contig139
SA-2129.1	. SA-2130.1	SA-2131.1	SA-2132.1	SA-2133.1	SA-2134.2
SeqID 1172	SeqID 1173	SeqID 1174	SeqID 1175	SeqID 1176	SeqID 1177
Sed	Sec	Seq	Sec	Sec	Sec



SeqID 1186	SA-2145.1	Contig114 (17361-18035 m)	64	Identities = 98/223 (43%), Positives = 145/223 (64%), Gaps = 5/223 (2%) dbj BAB04091:1 -(AP001508) two-component response regulator [Bacillus halodurans] Length = 222
SeqID 1187	SA-2146.1	Contig114 (18391-18525 m)	69	Identities = 33/44 (75%), Positives = 39/44 (88%) splP23376 RL34_BACST 50S RIBOSOMAL PROTEIN L34 pir  C48396 ribosomal protein L34 - Bacillus stearothermophilus gb AAB20570.1  BstL34=50S ribosomal subunit protein [Bacillus stearothermophilus, Peptide, 44 aa] gb AAB21085.1  ribosomal protein L34 [Bacillus stearothermophilus, Peptide, 44 aa] prf  1718186C ribosomal protein L34 [Bacillus stearothermophilus] Length = 44
SeqID 1188	SA-2147.1	Contig114 (18709-20064 m)	99	Identities = 211/459 (45%), Positives = 308/459 (66%), Gaps = 7/459 (1%) dbj BAB07666.1  (AP001520) unknown conserved protein [Bacillus halodurans] Length = 460
SeqID 1189	SA-2148.1	Contig114 (20311-22038 m)	73	Identities = 343/568 (60%), Positives = 426/568 (74%), Gaps = 2/568 (0%) gb AAF37879.1 AF234619_2 (AF234619) OpuABC [Lactococcus lactis] Length = 573
SeqID 1190	SA-2149.1	Ccntig114 (22057-23280 m)	81	Identities = 274/402 (68%), Positives = 337/402 (83%) gb AAF37878.1 AF234619 1 (AF234619) OpuAA [Lactococcus lactis] Length = 408
SeqID 1191	SA-2156.2	Contig98 (4979-5518 m)	. 62	Identities = 90/175 (51%); Positives = 118/175 (67%), Gaps = 2/175 (1%) splP36264 NUSG_STACA TRANSCRIPTION ANTITERMINATION PROTEIN NUSG pir  S38870 transcription antitermination factor nusG - Staphylococcus carnosus emb CAA53738.1  (X76134) nusG [Staphylococcus carnosus]
SeqID 1192	SA-2157.1	Contig98 (3727-4932 p)	38	Identities = 98/259 (37%), Positives = 155/259 (59%), Gaps = 10/259 (3%) gb AAF28363.1 AF224467_2 (AF224467) putative glycosyl transferase [Haemophilus ducreyi]



SeqID 1193 -SA-2158.1	SA-2158.1	Contig98 (2466-3662 p)	37 —	259 (5 AF224
				giycosyi transferase [riaemophilus ducieyi] Length = 209
;	:		:	Identities = 75/260 (28%), Positives = 123/260 (46%), Gaps = 22/260 (8%) sp P39407 YJJU_ECOLI HYPOTHETICAL 39.8 KD PROTEIN IN OSMY-DEOC INTERGENIC REGION (0357)
SeqID 1194	SA-2159.1	Contig98 (1416-2225·m)	43	mY-de 97273. 0.1  (A
. —				hypothetical protein [Escherichia coli K12] Length = 35/
SeqID 1195	SA-216.1	Contig136 (48774-49622 p)	45	Identities = 84/265 (31%), Positives = 133/265 (49%), Gaps = 14/265 (5%) gb AAB52383.1  (U36837) AbiEii [Lactococcus lactis] Length = 298
SeqID 1196	SA-2160.1	Contig98 (126-1373 p)	25	Identities = 139/391 (35%) Positives = 221/391 (55%), Gaps = 4/391 (1%) spiP71369jYB04_HAEIN HYPOTHETICAL METABOLITE TRANSPORT PROTEIN HI1104 pirjlC64167 hypothetical protein HI1104 - Haemophilus Influenzae (strain Rd
				KWZU) gbjAACZZ759.1j (U3Z79U) transporter protein [Haemophilus influenzae Rd] Length = 407
SeqID 1197	SA-2161.1	Contig81 (8-1291 m)	#	Identities = 35/101 (34%), Positives = 52/101 (50%), Gaps = 1/101 (0%) pdb 1BU  C Chain C, Structure Of The Ternary Microplasmin-Staphylokinase-
	-			Proteinase-Cofactor-Substrate Complex In Action Length = 128
SeqID 1198	SA-2162.1	Contig81 (1758-2456 p)	No Hits found	



S = 0 KD S = 1 KD S =	s = lyde }	TER llus ter ive cine	Gaps = in betL ttaine Length =
%), Gap AL 36.5 ir  E645 region 0007) y nypothe iH gene	(66%), Gaps ic semialdehy Length = 458	7178 (71%) TRANSPOR r opuD - Bacil aine transpor 108220) putat (Z99119) gly	%), Gap otein be s betain Leng
31 (519 HETIC SION p ergenic 11 (U00 5) off, b 3BD ye	55 (669 Sinic se Leng	27/178 E TRAI er opul etaine t -00822 1j (299 Lengt	24 (72% sport pr togene glycine es]
= 172/331 (5: HYPOTHET! NIC REGION P-nfo intergen A60511.1  (U( KE000305) orf,  2014253BD y  Length = 349	= 305/4 13) suca lis}	/es = 12 SETAIN ansport ycine b %.1  (Al %14985.	itives = 236/324 (72%) betaine transport pro monocytogenes I (AF102174) glycine i monocytogenes]
ositives ECOLI ERGEN in lysP gb AA gb AA 19.1  (AI	sitives F1025	Position (CINE Estaine transpose) global global global global mbjcAE subtilles subtilles	ositives ne betai _1 (AF1 mono
dentities_= 102/331 (30%), Positives = 172/331 (51%), Gaps = 6/331 (7%) splP33019 YEIH_ECOLI HYPOTHETICAL 36.9 KI PROTEIN IN LYSP-NFO INTERGENIC REGION pir  E64984 hypothetical 36.9 kD protein in lysP-nfo intergenic region - Escherichia coli (strain K-12) gb AAA60511.1  (U00007) yeiH Escherichia coli] gb AAC75219.1  (AE000305) orf, hypothetica protein [Escherichia coli K12] prf  2014253BD yeiH gene   Escherichia coli}	antities = 229/455 (50%), Positives = 3 55 (1%) gb AAD19405.1  (AF102543) dehydrogenase [Zymomonas mobilis]	Identities = 93/178 (52%), Positives = 127/178 (71%) 54417 OPUD_BACSU GLYCINE BETAINE TRANSPO UD pir  G69670 glycine betaine transporter opuD - Barbilis gb AAC4368.1  (U50082) glycine betaine transporter [Bacillus subtilis] gb AAC00408.1  (AF008220) put sporter [Bacillus subtilis] emb CAB14985.1  (Z99119) gbetaine transporter [Bacillus subtilis]	dentitles = 183/324 (56%), Positives = 236/324 (72%), Gaps 1/324 (0%) pir  T48645 glycine betaine transport protein betL [validated] - Listeria monocytogenes gb AAD30266.1 AF102174_1 (AF102174) glycine betaine ansporter BetL [Listeria monocytogenes] Length
/331 (3 P33019 YSP-N 36.9 kD 36.9 kD il (strai il gb[A herichia	/455 (5 AD194 ise [Zyr	= 93/17( D_BAC 9670 gly 544368 : subtilis illus sub	/324 (5   T4864 ated] -    6.1 AF
s = 102 7%) spl 3N IN L Sin IN L Shetical ichia col chia col sin [Escl	s = 229 %) gb A drogene	intities = 7/OPUI 7/OPUI gbl/A66 gbl/A66 Bacillus er [Baci	s = 183 (0%) pir [valid AD3026 ter BetL
Identities = 102/331 (30%), Positives = 172/331 (51%), Gaps = 26/331 (7%) splP33019 YEIH_ECOLI HYPOTHETICAL 36.9 KD PROTEIN IN LYSP-NFO INTERGENIC REGION pir  E64984 hypothetical 36.9 kD protein in lysP-nfo intergenic region - Escherichia coli (strain K-12) gb AAA60511.1  (U00007) yeil+ [Escherichia coli] gb AAC75219.1  (AE000305) orf, hypothetical protein [Escherichia coli K12] prf  2014253BD yeil+ gene   [Escherichia coli]	Identities = 229/455 (50%), Positives = 305/455 (66%), Gaps = 5/455 (1%) gb AAD19405.1  (AF102543) succinic semialdehyde dehydrogenase [Zymomonas mobilis]	Identities = 93/178 (52%), Positives = 127/178 (71%) splP54417(OPUD_BACSU GLYCINE BETAINE TRANSPORTER OPUD pirl G69670 glycine betaine transporter opuD - Bacillus subtilis gb AAC44368.1  (U50082) glycine betaine transporter OpuD [Bacillus subtilis] gb AAC00408.1  (AF008220) putative transporter [Bacillus subtilis] emb CAB14985.1  (Z99119) glycine betaine transporter [Bacillus subtilis]	Identities = 183/324 (56%), Positives = 236/324 (72%), Gaps = 1/324 (0%) pir  T48645 glycine betaine transport protein betL [validated] - Listeria monocytogenes gb AAD30266.1 AF102174_1 (AF102174) glycine betaine transporter BetL [Listeria monocytogenes] Length = 507
		s ∓	
. 4	63		02
			·
(d	1 p)	(d 6	1p)
(2627-3571 p)	(3645-5021 p)	(5164-5709 p)	(5710-6711 p)
. Contig81	Contig81	Contig81	Contig81
<del>-</del>	-		-
SA-2163.1	\$A-2165.1	SA-2166.1	SA-2167.1
SeqID 1199	SeqiD 1200	SeqID 1201	SeqID 1202
ა 	Sec	S	ŏ



SeqID 1203	SA-2168.1	Contig81 (6734-7219 m)		Identities = 68/152 (44%), Positives = 94/152 (61%), Gaps = 3/152 (1%) pir[IA70081 conserved hypothetical protein yxkH - Bacillus subtilis dbj[BAA11724.1 (D83026) homologous to SwissProt:YADE_ECOL!; hypothetical [Bacillus subtilis] emb[CAB15906.1 (Z99123) similar to hypothetical proteins [Bacillus subtilis]
SeqID 1204	SA-2169.1	Contig81 (7060-7620 m)	29	Identities = 38/122 (31%), Positives = 56/122 (45%), Gaps = 17/122 (13%) pir]A70081 conserved hypothetical protein yxkH - Bacillus subtilis dbj BAA11724.1  (D83026) homologous to SwissProt:YADE_ECOLI; hypothetical [Bacillus subtilis] emb CAB15906.1  (Z99123) similar to hypothetical proteins [Bacillus subtilis]
SeqID 1205	SA-217.1	Contig136 (48187-48777 p)	44	Identities = 53/219 (24%), Positives = 93/219 (42%), Gaps = 30/219 (13%) gb AAB52382.1[ (U36837) AbiEi [Lactococcus lactis] Length = 287
SeqID 1206	SA-2170.1	Contig§1 (7797-8006 p)	99	Identities = 35/69 (50%), Positives = 50/69 (71%) splP52985 DHOM_LACLA HOMOSERINE DEHYDROGENASE (HDH) pir JUC6049 homoserine dehydrogenase (EC 1.1.1.3) - Lactococcus lactis emb CAA65713.1  (X96988) hom [Lactococcus lactis] Length = 428
SeqID 1207	SA-2172.1	Contig80 (5804-7288 p)	61	Identities = 220/470 (46%), Positives = 311/470 (65%), Gaps = 16/470 (3%) pir  A82294 probable carbon starvation protein A VC0687 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF93852.1  (AE004154) carbon starvation protein A, putative [Vibrio cholerae]
SeqID 1208	SA-2173.1	Contig80 (4914-5648 p)	99	Identities = 93/245 (37%), Positives = 150/245 (60%), Gaps = 3/245 (1%) gb AAB48183.1  (L42945) lytR (Staphylococcus aureus)
SeqID 1209	SA-2174.1	Contig80 (3163-4902 p)	29	Identities = 265/582 (45%), Positives = 394/582 (67%), Gaps = 2/582 (0%) gb/AAB48182.1  (L42945) lytS (Staphylococcus aureus) Length = 584
SeqID 1210	SA-2175.1	Contig80 (2618-2782 p)	No Hits found	



				Identities = 283/489 (57%), Positives = 361/489 (72%), Gaps = 14/489 (2%) gb AAD20136 1  (AF091502) autoaggregation-mediating protein [Lactobacillus reuteri] Length = 497	Identities = 89/237 (37%), Positives = 132/237 (55%), Gaps = 3/237 (1%) pir[ C81348 probable periplasmic protein Cj0771c [imported] - Campylobacter jejuni (strain NCTC 11168) emb[CAB73036.1] (AL139076) putative periplasmic protein [Campylobacter jejuni]	Identities = 80/206 (38%), Positives = 127/206 (60%), Gaps = 4/206 (1%) pirl[D82957 probable permease of ABC transporter PA5504 [imported] - Pseudomonas aeruginosa (strain PAO1) gblAAG08889 11AE004963_2 (AE004963) probable permease of ABC transporter [Pseudomonas aeruginosa] Length = 225	Identities = 84/221 (38%), Positives = 138/221 (62%), Gaps = 5/221 (2%) sp P44785 ABC_HAEIN ATP-BINDING PROTEIN ABC pir  C64082 ATP-binding protein homolog H10621 - Haemophilus influenzae (strain Rd KW20) gb AAC22280.1  (U32744) ABC transporter, ATP-binding protein [Haemophilus influenzae Rd] Length = 345	Identities = 74/125 (59%), Positives = 92/125 (73%) pir  A69854 hypothetical protein yjqA - Bacillus subtilis emb CAB13104.1  (299110) yjqA [Bacillus subtilis] gb AAB87515.1  (AF034138) unknown [Bacillus subtilis] Length = 125
No Hits found	No Hits found	No Hits found	No Hits found	99		S S	δ	69
Contig80 (2239-2418 p)	Contig80 (1241-1618 p)	Contig80 (529-852 p)	Contig80 (2-181 p)	Contig129 (22347-23933 m)	Contig129 (24168-24998 m)	Contig129 (25014-25676 m)	Contig129 (25669-26403 m)	Contig129 (26524-26904 m)
SA-2177.2	SA-2178.1	SA-2180.2	SA-2182.2	SA-2184.1	SA-2185.1	SA-2186.1	SA-2187.1	SA-2188.1
SeqID 1211	SeqID 1212	SeqID 1213	SeqID 1214	SeqID 1215	SeqID 1216	SeqID 1217	SeqlD 1218	SeqID 1219



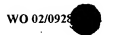
S	SA-2190.1	Contig129 (26990-28534 m)	75	Identities = 311/518 (60%), Positives = 393/518 (75%), Gaps = 9/518 (1%) spjO86490 RF3_STAAU PEPTIDE CHAIN RELEASE FACTOR 3 (RF-3) emb CAA74739.1  (Y14370) peptide chain release factor 3 [Staphylococcus aureus]
SA-2192.2	192.2	Contig129 (28711-30249 m)	56	Identities = 92/358 (25%), Positives = 142/358 (38%), Gaps = 55/358 (15%) gb AAG54632.1 AE005207_7 (AE005207) putative adhesin [Escherichia coli O157:H7] Length = 1417
SA-2	SA-2195.2	Contig109 (153-1748 m)	99	Identities = 243/373 (65%), Positives = 302/373 (80%), Gaps = 1/373 (0%) splP34001 YWAP_STRMU HYPOTHETICAL PROTEIN IN WAPA 3 REGION pir  S06993 hypothetical protein (wapA 3 region) - Streptococcus mutans (fragment) gb AAA88609.1  (M37842) unknown protein [Streptococcus mutans] Length = 373
SA-2	SA-2196.1	Contig109 (1867-3537 m)	87	Identities = 432/556 (77%), Positives = 492/556 (87%) sp Q59925 FTHS_STRMU FORMATETETRAHYDROFOLATE LIGASE (FORMYLTETRAHYDROFOLATE SYNTHETASE) (FHS) (FTHFS) gb AAB49329.1  (U39612) formyl-tetrahydrofolate synthetase [Streptococcus mutans] Length = 556
SA-S	SA-2197.1	Contig109 (3626-4645 m)	09	Identities = 131/331 (39%), Positives = 207/331 (61%), Gaps = 5/331 (1%) plr  G69830 lipoate-protein ligase homolog yhfJ - Bacillus subtilis emb CAA74531.1  (Y14083) hypothetical protein [Bacillus subtilis] emb CAB12865.1  (Z99109) similar to lipoate-protein ligase [Bacillus subtilis]
SA-	SA-2198.2	Contig109 (4672-5550 m)	No Hits found	
S, A	SA-22.1	Contig137 (22578-24410 p)	52	Identities = 228/571 (39%), Positives = 347/571 (59%), Gaps = 24/571 (4%) piral probable ATP-dependent proteinase ATP-binding protein NMA1045 [imported] - Neisseria meningitidis (group A strain Z2491) emb[CAB84311.1] (AL162755) putative ATP-dependent protease ATP-binding protein [Neisseria meningitidis] Length = 759



SeqID: 1227:-	SA-220.1	Contig136 (43070-47860 m) -	. 36.	Identities = 420/1385 (30%), Positives = 606/1385 (43%), Gaps = 308/1385 (22%) gb AAC44100.1  (U40026) SspB-precursor [Streptococcus gordonii] Length = 1500
SeqID 1228	SA-2200.2	Contig78 (4809-6092 p)		Identities = 365/427 (85%), Positives = 404/427 (94%)   sp O85730 TIG_STRPY TRIGGER FACTOR (TF)   gb AAC82391.1  (AF073922) RopA [Streptococcus pyogenes]   Length = 427
SeqID 1229	SA-2201.1	Contig78 (3810-4652 m)	. 52	Identities = 105/261 (40%), Positives = 150/261 (57%), Gaps = 2/261 (0%) dbjjBAB06385.1 (AP001516) unknown conserved protein [Bacillus halodurans]
SeqiD 1230	SA-2202.1	Contig78 (3204-3773 p)	89	Identities = 91/176 (51%), Positives = 115/176 (64%) sp P39157 YWLG_BACSU HYPOTHETICAL 19.4 KD PROTEIN IN SPOIIR-GLYC INTERGENIC REGION pir  140482 hypothetical protein ywlG - Bacillus subtilis emb CAA86109.1  (Z38002) Unknown [Bacillus subtilis] emb CAB15708.1  (Z99122) alternate gene name: ipc-33d [Bacillus subtilis] prf  2108403H ipc-33d gene [Bacillus subtilis]
SeqID 1231	SA-2203.1	Contig78 (2743-3207 p)	. 49	Identities = 46/148 (31%), Positives = 78/148 (52%), Gaps = 9/148 (6%) pir  G75153 hypothetical protein PAB2090 - Pyrococcus abyssi (strain Orsay) emb CAB49310.1  (AJ248284) hypothetical protein [Pyrococcus abyssi] Length = 199
SeqID 1232	SA-2204.1	Contig78 (1975-2733 p)	28	Identities = 95/253 (37%), Positives = 150/253 (58%), Gaps = 13/253 (5%) splP39610[THID_BACSU PHOSPHOMETHYLPYRIMIDINE KINASE (HMP-PHOSPHATE KINASE) (HMP-P KINASE) pir  S39707 phosphomethylpyrimidine kinase thID - Bacillus subtills emb CAA51608.1  (X73124) ipa-52r [Bacillus subtilis] emb CAB15828.1  (X99123) phosphomethylpyrimidine kinase [Bacillus subtilis]



SeqID 1233       SA-2205.1       Contig78 (1236-2012 p)       56         SeqID 1234       SA-2206.1       Contig78 (2-1165 p)       62         SeqID 1235       SA-2207.2       Contig139 (46557-47384 p)       73         SeqID 1236       SA-2208.2       Contig139 (47371-47844 p)       50         SeqID 1237       SA-2210.1       Contig139 (47856-49514 p)       63	2/240 (0%) spl0929J0jTRUA_BACHD TRNA PSEUDOURIDINE SYNTHASE A (PSEUDOURIDYLATE SYNTHASE I) (PSEUDOURIDINE SYNTHASE I) (URACIL HYDROLYASE) pril T44415 pseudourdyla:e synthase I truA [imported] - Bacillus halodurans dbjlBAA75303 11 (AB017508) truA homologue (identity of 62 to 8 subtilis%) [Bacillus halodurans] dbjlBAB03886 11 (AP001507) tRNA pseudouridine synthase A (pseudourdylate synthase
SeqID 1238 SA-2212.1 Contig139 (49627-50463 p) 54	Identities = 93/277 (33%), Positives = 152/277 (54%), Gaps = 4/277 (1%) dbj BAB07346.1  (AP001519) unknown conserved protein [Bacillus halodurans] Length = 283



Identities = 95/157 (60%), Positives = 121/157 (76%) gb AA86383.1  (U23376) putative 20-kDa protein [Lactococcus lactis] Length = 183 Identities = 20/43 (46%), Positives = 29/43 (66%), Gaps = 1/43 (2%) gb AA619662.1  (AE005054) calcium-binding protein homology; Cbp [Halobacterium sp. NRC-1] Length = 385 Identities = 24/61 (39%), Positives = 37/61 (60%), Gaps = 1/61 (1%) gb AAB96651.1  (AF034574) putative cruciform DNA binding protein [Glomus versiforme] Length = 99 Identities = 83/153 (54%), Positives = 110/153 (71%) gb AAA86383.1  (U23376) putative 20-kDa protein [Lactococcus lactis] Length = 183	53 58 .	Contig77 (4217-4759 m) Contig136 (42878-43069 m) Contig77 (3961-4158 m) Contig77 (3388-3939 m)	SA-2219.1 SA-222.1 SA-2220.1	SeqID 1244 SeqID 1245 SeqID 1246 SeqID 1246
Identities = 95/157 (60%), Positives = 121/157 (76%) gb AAA86383.1  (U23376) putative 20-kDa protein [Lactococcus lactis] Length = 183	64		SA-2219.1	SeqiD 1244
Identities = 27/61 (44%), Positives = 45/61 (73%)   gb AAA86382.1  (U23376) putative 6-kDa protein [Lactococcus   lactis]	99	Contig77 (4815-5009 m)	SA-2217.1	SeqID 1243
	No Hits found	Contig77 (5019-5570 m)	SA-2216.1	SeqID 1242
Identities = 31/84 (36%), Positives = 51/84 (59%) pir]B69770 conserved hypothetical protein ydaS - Bacillus subtilis dbj[BAA19274.1  (AB001488) FUNCTION UNKNOWN. [Bacillus subtilis] emb CAB12244.1  (Z99106) similar to hypothetical proteins from B. subtilis [Bacillus subtilis] Length = 85	55	Contig77 (5615-5863 m)	SA-2215.1	SeqID 1241
Identities = 75/185 (40%), Positives = 116/185 (62%), Gaps = 3/185 (1%) emb CAA72097.1  (Y11213) hypothetical protein [Streptococcus thermophilus]	25	Contig139 (51270-51872 p)	SA-2214.2	SeqID 1240
Identities = 149/274 (54%), Positives = 208/274 (75%), Gaps = 9/274 (3%) emb CAA72096:1  (Y11213) hypothetical protein [Streptococcus thermophilus]	69	Contig139 (50531-51295 p)***	SA-2213.1	SeqID 1239



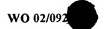
- <u>-</u>	•	<u> </u>	<u> </u>	Τ	<u> </u>	$\overline{}$	, O. <u>a</u>
Identities = 22/48 (45%), Positives = 35/48 (72%)  ref[NP_070072.1  A. fulgidus predicted coding region AF1244 [Archaeoglobus fulgidus] pir  C69405 hypothetical protein AF1244 - Archaeoglobus fulgidus gb AAB90005.1  (AE001018) A. fulgidus predicted coding region AF1244.  [Archaeoglobus fulgidus]	Identities = 67/262 (25%), Positives = 121/262 (45%), Gaps = 21/262 (8%) pir]B72352 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD35735.1 AE001738_15 (AE001738) conserved hypothetical protein [Thermotoga maritima] Length = 268	Identities = 67/218 (30%), Positives = 120/218 (54%), Gaps = 5/218 (2%) emb CAB40581.1  (AJ010128) DNA alkylation repair enzyme [Bacillus cereus] Length = 237	Identities = 119/163 (73%), Positives = 145/163 (88%) splP31308JTPX_STRSA PROBABLE THIOL PEROXIDASE pir  B43583 thioredoxin peroxidase (EC 1.11.1) - Streptococcus sanguis gb AAC98427.1  (M63481) 20-kDa protein [Streptococcus sanguinis] Length = 163		Identities = 39/135 (28%), Positives = 76/135 (55%), Gaps = 9/135 (6%) gb AAG09977.1 AF248038_6 (AF248038) GatA [Streptococcus agalactiae] Length = 149		Identities = 112/408 (27%), Positives = 197/408 (47%), Gaps = 21/408 (5%) splP39365 SGCC_ECOLI PUTATIVE PHOSPHOTRANSFERASE ENZYME II, C COMPONENT SGCC pir[ S56529 probable phosphotransferase enzyme II - Escherichia coli gb AAA97200.1  (U14003) ORF_f437 [Escherichia coli] gb AAC77260.1  (AE000501) putative PTS system enzyme IIC component [Escherichia coli K12]
15	94	52	85	No Hits found	46	No Hits found	40
Contig77 (2583-3224 m)	Contig77 (1814-2578 m)	Contig77 (1155-1814 m)	Contig77 (588-1082 m)	Contig77 (3-512 p)	Contig133 (18629-19093 m)	Contig133 (18322-18627 m)	Contig133 (16834-18282 m)
SA-2222.1	SA-2223.1	SA-2224.1	SA-2225.1	SA-2226.1	SA-2227.2	SA-2228.1	SA-2231.1
SeqID 1248	SeqID 1249	SeqID 1250	SeqID 1251	SeqID 1252	SeqID 1253	SeqID 1254	SeqID 1255



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Identities = 294/504 (58%), Positives = 380/504 (75%), Gaps = 10/504 (1%) splQ59959INANA_STRPN SIALIDASE A PRECURSOR (NEURAMINIDASE A) pir[IT30287 exo-alphasialidase (EC 3.2.1.18) - Streptococcus pneumoniae emb CAA51473.1  (X72967) neuraminidase (Streptococcus pneumoniae) Length = 1035	Identities = 57/146 (39%), Positives = 81/146 (55%), Gaps = 8/146 (5%) gb AAF18951.1 AF155805_5 (AF155805) Cps9H [Streptococcus suis]	Identities = 108/327 (33%), Positives = 170/327 (51%), Gaps = 22/327 (6%) emb CAC14890.1  (AJ295156) d-TDP-glucose dehydratase [Phragmites australis]	Identities = 77/231 (33%), Positives = 131/231 (56%), Gaps = 6/231 (2%) pir  S66119 conserved hypothetical protein yacM - Bacillus subtilis dbj  BAA05324.1  (D26185) unknown [Bacillus subtilis] emb  CAB11866.1  (299104) similar to hypothetical proteins [Bacillus subtilis] Length = 232	Identities = 85/274 (31%), Positives = 141/274 (51%), Gaps = 16/274 (5%) gb AAD37093.1 AF106539_2 (AF106539) LicD1 [Streptococcus pneumoniae] Length = 267		Identities = 27/109 (24%), Positives = 55/109 (49%) dbj BAA19645.1  (AB002668) unnamed protein product [Actinobacillus actinomycetemcomitans] Length = 126		Identities = 118/240 (49%), Positives = 152/240 (63%), Gaps = 1/240 (0%) gb[AAC35924.1] (AF071085) putative glycosyl transferase [Enterococcus faecalis] Length = 241
45	25	47	52	50	No Hits found	46	No Hits found	69
Contig133 (14146-16596 m)	Contig101 (6129-7064 p)	Contig101 (5072-6127 p)	Contig101 (4347-5069 p)	Contig101 (3523-4347 p)	Contig101 (1765-3498 p)	i	Contig136 (42343-42894 m)	Contig101 (691-1422 p)
SA-2232.2	SA-2233.2	SA-2235.1	SA-2236.1	SA-2237.1	SA-2238.1	SA-2239.1	SA-224.1	SA-2240.1
SeqID 1256	SeqID 1257	SeqID 1258	SeqID 1259	SeqID 1260	SeqID 1261	SeqID 1262	SeqID 1263	SeqID 1264



SeqID 1265	SA-2241.2	Contig101 (3-689 p)	75	Identities = 123/231 (53%), Positives = 174/231 (75%), Gaps = 7/231 (3%) pir  T00087 rhamnosyltransferase- Streptococcus mutans dbj  BAA32090.1  (AB010970) rhamnosyltransferase   [Streptococcus mutans]
SeqID 1266	SA-2242.2	Contig115 (41-406 p)	75	Identities = 77/118 (65%), Positives = 102/118 (86%) gb AAK04289.1 AE006256_11 (AE006256) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 122
SeqID 1267	SA-2244.2	Contig115 (406-2070 p)	78	Identitles = 354/539 (65%), Positives = 438/539 (80%), Gaps = 8/539 (1%) gb AAK04288.1 AE006256_10 (AE006256) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 535
SeqID 1268	SA-2245.1	Contig115 (2254-3102 p)	56	Identities = 127/275 (46%), Positives = 174/275 (63%), Gaps = 1/275 (0%) gb AAF68390.1 AF236374_1 (AF236374) hypersensitive-induced response protein [Zea mays] Length = 284
SeqID 1269	SA-2246.1	Contig115 (4134-4664 p)	31	Identities = 39/110 (35%), Positives = 55/110 (49%), Gaps = 3/110 (2%) pir  G72536 hypothetical protein APE1580 - Aeropyrum pernix (strain K1) dbj BAA80580.1  (AP000062) 114aa long hypothetical protein [Aeropyrum pernix] Length = 114
SeqID 1270	SA-2247.1	Contig115 (4184-4924 m)	71	Identities = 132/241 (54%), Positives = 178/241 (73%), Gaps = 1/241 (0%) pir  F81363 probable glutamine transport ATP-binding protein Cj0902 [imported] - Campylobacter jejuni (strain NCTC 11168) emb CAB73160.1  (AL139076) putative glutamine transport ATP-binding protein [Campylobacter jejuni]
SeqID 1271	SA-2248.2	Contig115 (4934-6484 m)	49	Identities = 147/534 (27%), Positives = 255/534 (47%), Gaps = 75/534 (14%) pir  S77250 hypothetical protein - Synechocystis sp. (strain PCC 6803) dbj BAA17584.1  (D90907) glutamine-binding periplasmic protein [Synechocystis sp.] Length = 530
SeqID 1272	SA-225.1	Contig136 (41699-42292 m)	No Hits found	



Saps = 3/95 : ABC oir  B69956 gH - Bacillus us subtilis] ubtilis] zmC~similar ; [Bacillus	80%) VSPORTER Shate ABC subtilis ubtilis]	%), Gaps = BLE ABC pir  B69956 igH - Bacillus us subtilis  zmC~similar zmC~similar   [Bacillus
Identities = 35/95 (36%), Positives = 56/95 (58%), Gaps = 3/95  TRANSPORTER PERMEASE PROTEIN YQGH piri[B69956 phosphate ABC transporter (permease) homolog yqgH - Bacillus subtilis dbj[BAA09582.1] (D58414) ORF72 [Bacillus subtilis] dbj[BAA12511.1] (D84432) YqgH [Bacillus subtilis] emb[CAB14428.1] (299116) alternate gene name: yzmC~similar to phosphate ABC  transporter (permease) [Bacillus subtilis] Length = 309	Identities = 35/54 (64%), Positives = 44/54 (80%)  sp P46339 YQGH_BACSU PROBABLE ABC TRANSPORTER PERMEASE PROTEIN YQGH pir  B69956 phosphate ABC transporter (permease) homolog yqgH - Bacillus subtilis dbj BAA09582.1  (D58414) ORF72 [Bacillus subtilis] dbj BAA12511.1  (D84432) YqgH [Bacillus subtilis] emb CAB14428.1  (299116) alternate gene name: yzmC-similar to phosphate ABC transporter (permease) [Bacillus subtilis] Length = 309	Identities = 78/161 (48%), Positives = 113/161 (69%), Gaps = 1/161 (0%) sp P46339 YQGH_BACSU PROBABLE ABC TRANSPORTER PERMEASE PROTEIN YQGH pir  B69956 phosphate ABC transporter (permease) homolog yqgH - Bacillus subtilis dbj BAA09582.1  (D58414) ORF72 [Bacillus subtilis] dbj BAA12511.1  (D84432) YqgH [Bacillus subtilis] emb CAB14428.1  (Z99116) alternate gene name: yzmC~similar to phosphate ABC transporter (permease) [Bacillus subtilis] Length = 309
Pho Pho	sp tra	b hq s
47	45	62
i (9163-9489 p)	t (9422-9700 p)	(9663-10082 p)
- Contig104	Contig 104	Contig 104
SA-2251.1.	SA-2252.1	SA-2253.1
SeqID 1273	SeqID 1274	SeqiD 1275



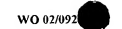
ooktek subtilis subtilis tilis] is] ioD~similar sacillus	32%) ATE 564426 n - J67544) pstB)	75%) 3C IID69956 og yggJ - DRF74 3acillus name: P-binding	76%) transport lactis]
Identities = 157/294 (53%), Positives = 225/294 (76%) sp P46340 YQGI_BACSU PROBABLE ABC TRANSPORTER PERMEASE PROTEIN YQGI pir  C69956 phosphate ABC transporter (permease) homolog yqgl - Bacillus subtilis dbj BAA09583.1  (D58414) ORF73 [Bacillus subtilis] dbj BAA12512.1  (D84432) Yqgl [Bacillus subtilis] emb CAB14427.1  (Z99116) alternate gene name: yzmD~similar to phosphate ABC transporter (permease) [Bacillus subtilis] . Length = 294	Identities = 154/247 (62%), Positives = 204/247 (82%) sp[Q58418 PSTB_METJA PROBABLE PHOSPHATE TRANSPORT ATP-BINDING PROTEIN PSTB pir  C64426 phosphate transport system ATP-binding protein - Methanococcus jannaschii gb AAB99016.1  (U67544) phosphate specific transport complex component (pstB) [Methanococcus jannaschii] Length = 252	Identities = 148/248 (59%), Positives = 189/248 (75%) splP46341 YQGJ_BACSU HYPOTHETICAL ABC TRANSPORTER ATP-BINDING PROTEIN YQGJ pir ID69956 phosphate ABC transporter (ATP-binding pro) homolog yqgJ-Bacillus subtilis dbj BAA09584.1  (D58414) ORF74 [Bacillus subtilis] dbj BAA12513.1  (D84432) YqgJ [Bacillus subtilis] emb CAB14426.1  (Z99116) alternate gene name: yzmE~similar to phosphate ABC transporter (ATP-binding protein) [Bacillus subtilis]	Identities = 116/217 (53%), Positives = 167/217 (76%) gb AAK05813.1 AE006402_1 (AE006402) phosphate transport system regulator [Lactococcus lactis subsp. lactis] Length = 217
74	74	. 47	72
Contig104 (1007210959 p)	Contig104 (10971-11774 p)	Contig104 (11786-12544 p)	Contig104 (12578-13231 p)
SA-2254.1	SA-2255.1	SA-2256.1	SA-2258.2
SeqiD 1276	SeqID 1277	SeqiD 1278	SeqID 1279



SeqID 1280	SA-2259.1	Contig76 (2249-4879 p)		Identities = 107/531 (20%), Positives = 224/531 (42%), Gaps = 62/531 (11%) ref[NP_070647.1] A. fulgidus predicted coding region AF1820 [Archaeoglobus fulgidus] pir[IC69477 hypothetical protein AF1820 - Archaeoglobus fulgidus gb[AAB89436.1] (AE000977) A. fulgidus predicted coding region AF1820 [Archaeoglobus fulgidus]
SeqID 1281	SA-2260.1	Contig76 (1536-2237 p)	70	Identities = 112/230 (48%), Positives = 167/230 (71%) ref[NP_070646.1  ABC transporter, ATP-binding protein [Archaeoglobus fulgidus] pir  B69477 ABC transporter, ATP-binding protein homolog - Archaeoglobus fulgidus gb AAB89431.1  (AE000977) ABC transporter, ATP-binding protein [Archaeoglobus fulgidus]
SeqID 1282	SA-2261.1	Contig76 (158-1399 p)	76	Identities = 278/469 (59%), Positives = 355/469 (75%), Gaps = 10/469 (2%) splQ9KA23 TOP1_BACHD DNA TOPOISOMERASE I (OMEGA-PROTEIN) (RELAXING ENZYME) (UNTWISTING ENZYME) (SWIVELASE) dbj BAB06186.1  (AP001515) DNA topoisomerase I [Bacillus halodurans]
SeqID 1283	SA-2263.1	Contig 127 (39359-39559 p)	No Hits found	
SeqiD 1284	SA-2264.2	Contig127 (39121-39972 m)	64	Identities = 126/284 (44%), Positives = 185/284 (64%) splP37550jiSPE_BACSU 4-DIPHOSPHOCYTIDYL-2-C-METHYL. D-ERYTHRITOL KINASE (CMK) (4-(CYTIDINE-5 - DIPHOSPHO)-2-C-METHYL-D-ERYTHRITOL KINASE) pirjS66075 conserved hypothetical protein yabH - Bacillus subtilis dbjjBAA05281.1  (D26185) unknown [Bacillus subtilis] embjCAB11822.1  (Z99104) similar to hypothetical proteins [Bacillus subtilis] Length = 289
SeqID 1285	. SA-2265.2	Contig127 (38592-39035 m)	73	Identities = 77/146 (52%), Positives = 117/146 (79%) pir  T46753 repressor protein adcR [imported] - Streptococcus pneumoniae emb CAA96184.1  (Z71552) AdcR protein [Streptococcus pneumoniae]



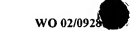
SeqID 1286	SA-2266.2	Contig127 (37879-38589 m)	84	Identities = 182/231 (78%), Positives = 206/231 (88%) pir  T46754 AdcC-protein [imported] - Streptococcus pneumoniae emb CAA96186.1  (Z71552) AdcC protein [Streptococcus pneumoniae] Length = 234
SeqID 1287	SA-2267.1	Contig127 (37077-37889 m)	11	Identities = 197/263 (74%), Positives = 236/263 (88%) pir  T46755 membrane protein adcB [imported] - Streptococcus pneumoniae emb CAA96187.1  (Z71552) AdcB protein [Streptococcus pneumoniae] Length = 268
SeqID 1288	SA-2268.2	Contig127 (35834-36865 p)	41	Identities = 116/216 (53%), Positives = 150/216 (68%), Gaps = 9/216 (4%) gb AAK04254.1 AE006253_5 (AE006253) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 263
SeqiD 1289	SA-2269.1	Contig134 (81376-82521 m)	29	Identities = 169/374 (45%), Positives = 235/374 (62%), Gaps = 16/374 (4%) splO06005 AAPA_BACSU AMINO ACID PERMEASE AAPA pir B69580 amino acid permease aapA - Bacillus subtilis emb CAA63459.1  (X92868) amino acid permease [Bacillus subtilis] emb CAB14651.1  (Z99117) amino acid permease [Bacillus subtilis]
SeqID 1290	SA-2271.1	Contig134 (80825-81292 p)	98	Identities = 121/155 (78%), Positives = 139/155 (89%)   gb AAC23745.1  (AF052209) VacB homolog [Streptococcus   pneumoniae]
SeqID 1291	SA-2272.2	Contig134 (78417-80822 p)	61	Identities = 350/815 (42%), Positives = 501/815 (60%), Gaps = 49/815 (6%) spl032231fRNR_BACSU RIBONUCLEASE R (RNASE R) (VACB PROTEIN HOMOLOG) pirl[G70027 conserved hypothetical protein yvaJ - Bacillus subtilis emb CAB15366.1  (299121) similar to hypothetical proteins [Bacillus subtilis]
SeqID 1292	SA-2274.1	Contig123 (1-786 m)	92	Identities = 169/260 (65%), Positives = 210/260 (80%), Gaps = 1/260 (0%) gb AAF86640.1 AF162694_1 (AF162694) ABC transporter [Enterococcus gallinarum] Length = 269



Identities = 114/285 (40%), Positives = 175/285 (61%), Gaps = 3/285 (1%) pirijF83165 probable permease of ABC transporter PA3837 [imported] - Pseudomonas aeruginosa (strain PAO1) gbjAAG07224 1JAE004801_2 (AE004801) probable permease of ABC transporter [Pseudomonas aeruginosa] Length = 296	Identities = 103/283 (36%). Positives = 165/283 (57%), Gaps = 1/283 (0%) pir[[E83165 hypothetical protein PA3836 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG07223.1 AE004801_1 (AE004801) hypothetical protein [Pseudomonas aeruginosa] Length = 325	Identities = 89/261 (34%), Positives = 151/261 (57%), Gaps = 4/261 (1%) sp P4447 Y003_HAEIN PROTEIN HI0003 pir[  64139 hypothetical protein HI0003 - Haemophilus influenzae (strain Rd KW20) gb AAC21682.1  (U32686) conserved hypothetical protein [Haemophilus influenzae Rd] Length = 262	Identities = 86/275 (31%), Positives = 143/275 (51%), Gaps = 6/275 (2%) dbj BAB07024 1  (AP001518) unknown conserved protein [Bacillus halodurans] Length = 286	Identities = 97/240 (40%), Positives = 151/240 (62%), Gaps = 1/240 (0%) gb AAD24446.1 AF118389_3 (AF118389) unknown [Streptococcus suis] Length = 244	No Hits found No Hits found	Identities = 437/522 (83%), Positives = 484/522 (92%), Gaps = 7/522 (1%) splQ54431 SR54_STRMU SIGNAL RECOGNITION PARTICLE PROTEIN (FIFTY-FOUR HOMOLOG) gb AAB48050.1 AAB48050 (U88582) Ffh [Streptococcus mutans] Length = 516	Identities = 95/110 (86%), Positives = 103/110 (93%)   84   gb AAB48049.1 AAB48049 (U88582) YIXM [Streptococcus
Contig123 (791-1687 m)	Contig123 (1703-2647 m)	Contig123 (3130-3939 m)	Contig123 (4113-4973 p)	Contig123 (5016-5747 p)	Contig104 (20416-20709 p) Contig104 (20191-20379 p)	1	Contig104 (18154-18486 p)
SA-2276.1	SA-2278.1	SA-2279.1	SA-2280.1	SA-2281.2	SA-2282.1 SA-2283.1	SA-2285.1	SA-2287.1
SeqID 1293	SeqID 1294	SeqID 1295	SeqID 1296	SeqID 1297	SeqID 1298 SeqID 1299	SeqID 1300	SeqID 1301

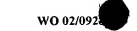


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Identities = 219/420 (52%), Positives = 306/420 (72%), Gaps = 4/420-(0%) splQ54955 CIAH_STRPN SENSOR PROTEIN CIAH-pir  S49545 histidine kinase - Streptococcus pneumoniae emb CAA54466.1  (X77249) histidine kinase [Streptococcus pneumoniae] emb CAB54565.1  (AJ005926) histidine kinase [Streptococcus pneumoniae]	Identities = 152/245 (62%), Positives = 193/245 (78%), Gaps = 1/245 (0%) gb AAK06239.1 AE006442_6 (AE006442) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 250	Identities = 66/200 (33%), Positives = 111/200 (55%), Gaps = 10/200 (5%) pir  E69826 probable 1-acylglycerol-3-phosphate O-acyltransferase (EC 2.3.1.51) yhdO - Bacillus subtilis emb CAA74499.1  (Y14082) hypothetical protein [Bacillus subtilis] emb CAB12793.1  (299109) similar to 1-acylglycerol-3-phosphate O-acyltransferase [Bacillus subtilis] Length = 199	Identities = 96/217 (44%), Positives = 138/217 (63%), Gaps = 4/217 (1%) gbjAAC23741.1  (AF052208) competence protein [Streptococcus pneumonlae] Length = 216	Identities = 120/286 (41%), Positives = 180/286 (61%), Gaps = 1/286 (0%) gb AAC23742.1  (AF052208) competence protein [Streptococcus pneumoniae]	Identities = 205/419 (48%), Positives = 298/419 (70%), Gaps = 2/419 (0%) gb AAC23742.1  (AF052208) competence protein [Streptococcus pneumoniae] Length = 753	Identities = 120/267 (44%), Positives = 177/267 (65%), Gaps = 6/267 (2%) gblAAK04342.1JAE006262_1 (AE006262) conserved hypothetical protein [Lactococcus lactis subsp. lactis]  Length = 270
69	73	4	09	09	64	61
Contig104 (16752-18065 p)	Contig129 (21139-21903 p)	Contig129 (20264-21004 m)	Contig129 (19511-20164 m)	Contig129 (18655-19527 m)	Contig129 (17289-18554 m)	Contig129 (16354-17163 m)
SA-2288.2	SA-2289.2	SA-2291.1	SA-2292.1	SA-2293.1	SA-2294.1	SA-2295.2
SeqID 1302	SeqID 1303	SeqID 1304	SeqID 1305	SeqID 1306	SeqID 1307	SeqID 1308



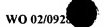
SeqID 1309	SA-2296.2	Contig101 (10807-11655 p)	57	Identities = 112/278 (40%), Positives = 163/278 (58%), Gaps = 2/278 (0%) emb CAB52237.1  (Z98171) EpsQ protein [Streptococcus thermophilus]
SeqID 1310	SA-2297.1	Contig101 (11645-12784 p)	. 51	Identities = 108/395 (27%), Positives = 196/395 (49%), Gaps = 28/395 (7%) pir  D64433 hypothetical protein MJ1069 - Methanococcus jannaschii gb AAB99071.1  (U67549) galactosyltransferase isolog [Methanococcus jannaschii] Length = 392
SeqID 1311	SA-2298.1	Contig101 (12836-14365 m)	54	Identities = 172/492 (34%), Positives = 289/492 (57%), Gaps = 2/492 (0%) gb AAC97147.1  (U49397) Nra [Streptococcus pyogenes] Length = 511
SeqID 1312	SA-23.1	Contig137 (20828-22576 p)	53	dentities = 218/608 (35%), Positives = 323/608 (52%), Gaps = 52/608 (8%) gb AAC38606.1  (AF007787) type I topoisomerase [Enterococcus faecalis]
SeqID 1313	SA-230.1	Contig136 (35495-41695 m)	29	Identities = 373/1243 (30%), Positives = 614/1243 (49%), Gaps = 97/1243 (7%) ref[NP_066674.1] similar to jhp0928 gene in Helicobacter pylori [Agrobacterium rhizogenes] dbj BAB16212.1  (AP002086) similar to jhp0928 gene in Helicobacter pylori [Agrobacterium rhizogenes] Length = 1693
SeqID 1314	SA-2300.3	Contig101 (14589-17354 p)	6	Identities = 58/176 (32%), Positives = 83/176 (46%), Gaps = 19/176 (10%) gb AAD33086.1 AF071083_1 (AF071083) fibronectin-binding protein I [Streptococcus pyogenes] Length = 1161
SeqID 1315	SA-2302.2	Contig79 (4055-5029 p)		Identities = 138/315 (43%), Positives = 222/315 (69%), Gaps = 6/315 (1%) pir[]C69763 ferrichrome ABC transporter (permease) homolog yclO - Bacillus subtilis dbj[BAA09013.1] (D50453) homologue of ferric anguibactin transport system permerase protein FatC of V. anguillarum [Bacillus subtilis] emb[CAB12189.1] (299106) similar to ferrichrome ABC transporter (permease) [Bacillus subtilis] Length = 315

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Identities = 149/304 (49%). Positives = 234/304 (76%) pir[]B69763 ferrichrome ABC transporter (permease) homolog yclN - Bacillus subtilis abj[BAA09012.1] (D50453) homologue of ferric anguibactin transport system permerase protein FatD of V. anguillarum [Bacillus subtilis] emb[CAB12188 1] (299106) similar to ferrichrome ABC transporter (permease) [Bacillus subtilis] Length = 316	Identities = 93/182 (51%), Positives = 125/182 (68%), Gaps = 2/182 (1%) dbjjBAB06720.1  (AP001517) maltose transacetylase (maltose O-acetyltransferase) [Bacillus halodurans] Length = 186	Identities = 128/249 (51%), Positives = 168/249 (67%) sp O31744 RNH2_BACSU RIBONUCLEASE HII (RNASE HII) pir  C69693 ribonuclease H rnh - Bacillus subtilis emb CAB13479.1  (299112) ribonuclease H [Bacillus subtilis] Length = 255	Identities = 141/281 (50%), Positives = 196/281 (69%), Gaps = 5/281 (1%) dbj BAA75361.1  (AB013365) YlqF [Bacillus halodurans] dbj BAB06195.1  (AP001515) unknown conserved protein [Bacillus halodurans] Length = 284	Identities = 61/135 (45%), Positives = 86/135 (63%), Gaps = 4/135 (2%) pir  A69760 conserved hypothetical protein yciB - Bacillus subtilis dbj BAA08969.1  (D50453) yciB [Bacillus subtilis] emb CAB12129.1  (299105) similar to hypothetical proteins [Bacillus subtilis] Length = 194	Identities = 85/336 (25%), Positives = 158/336 (46%), Gaps = 28/336 (8%) pir  E71665 bicyclomycin resistance protein (bcr1) RP603 - Rickettsia prowazekii emb CAA15047.1  (AJ235272) BICYCLOMYCIN RESISTANCE PROTEIN (bcr1) [Rickettsia prowazekii] Length = 407
. 67	63	64	99	62	40
Contig79 (3096-4058 p)	Contig79 (2309-2857 p)	Contig79 (1524-2288 p)	Contig79 (689-1540 p)	Contig79 (72-413 p)	Contig74 (5434-6534 m)
SA-2303.1	SA-2304.1	SA-2305.1	SA-2306.1	SA-2307.1	SA-2308.1
SeqID 1316.	SeqID 1317	SeqID 1318	SeqID 1319	SeqiD 1320	SeqID 1321



SeqID 1322	SA-2309.1	Contig74 (4435-5382 m)	53	Identities = 127/269 (47%), Positives = 177/269 (65%), Gaps = 8/269 (2%) gb AAC23746.1  (AF052209) competence protein [Streptococcus pneumoniae] Length = 266
SeqID 1323	SA-231.1	Contig136 (35124-35423 m)	No Hits found	
SeqID 1324	SA-2311.1	Contig74 (2614-4419 m)	88	Identities = 593/601 (98%), Positives = 597/601 (98%) splQ53778 PEPB_STRAG GROUP B OLIGOPEPTIDASE PEPB pir  T51748 thimet oligopeptidase (EC 3.4.24.15) PepB [validated] Streptococcus agalactiae gb AAC44215.1  (U49821) group B oligopeptidase PepB [Streptococcus agalactiae] Length = 601
SeqID 1325	SA-2312.1	Contig74 (1793-2419 m)	28	Identities = 39/117 (33%), Positives = 67/117 (56%), Gaps = 9/117 (7%) pir[ A57362 gyrb protein - Streptococcus pneumoniae (fragment) emb CAA58770.1  (X83917) orflgyrb [Streptococcus pneumoniae] emb CAA91552.1  (Z67740) unidentified [Streptococcus pneumoniae]
SeqID 1326	SA-2313.1	Contig74 (1012-1719 m)	69	Identities = 131/227 (57%), Positives = 169/227 (73%)   emb CAA68045.1  (X99710) methyltransferase [Lactococcus   lactis]
SeqID 1327	SA-2314.1	Contig74 (22-951 m)	8	Identities = 101/307 (32%), Positives = 151/307 (48%), Gaps = 17/307 (5%) splP15294 PRTM_LACLA PROTEASE MATURATION PROTEIN PRECURSOR pir  S08083 probable protein export protein prtM precursor - Lactococcus lactis subsp. cremoris (strain NCDO 763) plasmid pLP763 emb CAA32349.1  (X14130) ORF (AA 1 to 299) [Lactococcus lactis subsp. cremoris]
SeqID 1328	SA-2315.2	Contig123 (8457-10655 m)	82	Identities = 539/725 (74%), Positives = 616/725 (84%), Gaps = 7/725 (0%) gbjAAD00215.1j (U73336) anaerobic ribonucleotide reductase [Lactococcus lactis subsp. cremoris] Length = 747
SeqID 1329	SA-2316.1	Contig123 (8239-8382 m)	No Hits found	

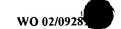
SeqID 1330	SA-2317.1	Contig123 (7294-8226 m)· -	46	Identities = 91/299 (30%), Positives = 147/299 (48%), Gaps = 7/299 (2%) emb CAB95794; 1]_(AL359949) putative oxidoreductase [Streptomyces coelicolor A3(2)]  301
SeqID 1331	SA-2318.1		40	Identities = 52/129 (40%), Positives.= 70/129 (53%), Gaps = .5/129 (3%) dbjjBAB04222.1  (AP001508) unknown conserved protein in others [Bacillus halodurans] Length = 174
SeqID 1332	SA-2319.2	Contig123 (6104-6721 m)	62	Identities = 152/198 (76%), Positives = 176/198 (88%) gb AAD00216.1  (U73336) anaerobic ribonucleotide reductase activator protein [Lactococcus lactis subsp. cremoris] Length = 199
SeqID 1333	SA-232.1	Contig136 (34814-35113 m)	No Hits found	
SeqID 1334	SA-2322.2	Contig118 (25862-27211 p)	76	Identities = 275/450 (61%), Positives = 347/450 (77%), Gaps = 1/450 (0%) dbj BAA76640.1  (AB019579) glutathione reductase (GR) [Streptococcus mutans] Length = 450
SeqID 1335	SA-2324.1	Contig118 (27254-27706 m)	90	Identities = 45/156 (28%), Positives = 79/156 (49%), Gaps = 3/156 (1%) gbjAAF87093.1JAF167576_1 (AF167576) secreted antigen SagBb [Enterococcus hirae] Length = 576
SeqID 1336	SA-2326.1	Contig118 (27953-29098 p)	09	Identities = 175/353 (49%), Positives = 234/353 (65%), Gaps = 1/353 (0%) splP31672 NIFS_LACDE NIFS PROTEIN HOMOLOG pir  S16047 nitrogenase cofactor synthesis protein nifS - Lactobacillus delbrueckii emb CAA43493.1  (X61190) nifS-like gene [Lactobacillus delbrueckii] Length = 355
SeqID 1337	SA-2327.2	Contig118 (29100-30314 p)	74	Identities = 264/385 (68%), Positives = 312/385 (80%)   gb AAK04477.1 AE006275_1 (AE006275) HYPOTHETICAL   PROTEIN [Lactococcus lactis subsp. lactis]
SeqID 1338	SA-2329.3	Contig118 (30416-31594 p)	28	Identities = 68/224 (30%), Positives = 117/224 (51%), Gaps = 10/224 (4%) ref[NP_053211.1  pXO2-56 [Bacillus anthracis] gb AAF13661.1 AF188935_59 (AF188935) pXO2-56 [Bacillus anthracis] Length = 411



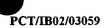
-			[s]	Ø	l	<b>£</b>	<sub>(O</sub>
	Identities = 260/293 (88%), Positives = 276/293 (93%)   dbj BAB16889 1 (AB050113) class-II aldolase [Streptococcus bovis]   Length = 293	Identities = 175/306 (57%), Positives = 220/306 (71%), Gaps = 3/306 (0%) splP14295 DHL2_LACCO L-2-HYDROXYISOCAPROATE DEHYDROGENASE (L-HICDH) pir  JQ0114 L-2-hydroxyisocaproate dehydrogenase (EC 1.1.1) Lactobacillus confusus gb AAA88213.1  (M31425) L-2-hydroxyisocaproate dehydrogenase [Weissella confusa] Length = 310	Identities = 45/62 (72%), Positives = 53/62 (84%) sp P37807 RL28_BACSU 50S RIBOSOMAL PROTEIN L28 pir  S39982 ribosomal protein L28 (rpmB) - Bacillus subtilis gb AAC36810.1  (L12244) ribosomal protein L28 (Bacillus subtilis] emb CAA74255.1  (Y13937) putative RpmB protein [Bacillus subtilis] emb CAB13455.1  (Z99112) ribosomal protein L28 [Bacillus subtilis] Length = 62	Identities = 65/236 (27%), Positives = 92/236 (38%), Gaps = 47/236 (19%) pir  S52348 hypothetical protein 2 - Lactobacillus leichmannii emb CAA57459.1  (X81869) orf2 [Lactobacillus leichmannii] Length = 507	Identities = 91/298 (30%), Positives = 155/298 (51%), Gaps = 13/298 (4%) gb AAK04857.1 AE006309_6 (AE006309) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 432	Identities = 95/271 (35%), Positives = 139/271 (51%), Gaps = 16/271 (5%) gb AAC13546.1  (AF019629) putative fimbria-associated protein [Actinomyces naeslundii] Length = 365	Identities = 62/245 (25%), Positives = 96/245 (38%), Gaps = 45/245 (18%) dbj BAB04080.1  (AP001508) unknown [Bacillus halodurans]
No Hits found	93	. 09	72	13	50	46	30
Contig136 (34011-34712 m)	Contig73 (1490-2371 m)	Contig73 (488-1405 p)	Contig73 (61-249 m)	. Contig101 (17522-19501 p)	Contig101 (19712-20491 p)	Contig101 (20491-21375 p)	Contig101 (21423-22298 p)
SA-233.1	SA-2330.1	SA-2331.1	SA-2332.1	SA-2334.1	SA-2335.2	SA-2336.1	SA-2337.1
SeqID 1339	SeqID 1340	SeqID 1341	SeqID 1342	SeqID 1343	SeqID 1344	SeqID 1345	SeqID 1346

		<b>-</b>				
Identities = 228/232 (98%), Positives = 229/232 (98%), Gaps = 3/232 (1%) dbj BAA82278.1  (AB028896) CpslaD [Streptococcus agalactiae]	Identities = 103/342 (30%), Positives = 155/342 (45%), Gaps = 50/342 (14%) ref NP_053232.1  pXO2-78 [Bacillus anthracis] gb AAF13682.1 AF188935_80 (AF188935) pXO2-78 [Bacillus anthracis] Length = 344	Identities = 448/449 (99%), Positives = 448/449 (99%) pir  T44643 galactosyl transferase cpsD [imported] - Streptococcus agalactiae gb AAD53066.1 AF163833_6 (AF163833) CpsE [Streptococcus agalactiae] Length = 449	Identities = 149/149 (100%), Positives = 149/149 (100%) pir  T44644 glycosyl transferase activity enhancer cpsG [imported] - Streptococcus agalactiae gb AAD53067.1 AF163833_7 (AF163833) CpsF [Streptococcus agalactiae] Length = 149	Identities = 155/157 (98%), Positives = 155/157 (98%) pir  T44645 glycosyl transferase cpsH [imported] - Streptococcus agalactiae gb AAD53068.1 AF163833_8 (AF163833) CpsG [Streptococcus agalactiae] Length = 157	Identities = 380/381 (99%), Positives = 380/381 (99%) pir  T44646 capsular polysaccharide repeating unit polymeras cpsl [imported] - Streptococcus agalactiae gb AAD53069.1 AF163833_9 (AF163833) CpsH [Streptococcus agalactiae] Length = 381	Identities = 318/322 (98%), Positives = 320/322 (98%) pir  T44647 glycosyl transferase cpsJ [imported] - Streptococcus agalactiae gb AAD53070.1 AF163833_10 (AF163833) CpsI [Streptococcus agalactiae] Length = 322
96	38	96	63	94	66	86
Contig135 (2828-3526 p)	Contig136 (32885-33970 m)	Contig135 (3539-4927 p)	Contig135 (4951-5400 p)	Contig135 (5400-5873 p)	Contig135 (5870-7015 p)	Contig135 (7012-7980 p)
SA-2339.2	SA-234.1	SA-2340.2	SA-2341.1	SA-2342.1	SA-2343.1	SA-2344.1
SeqID 1347	SeqID 1348	SeqID 1349	SeqID 1350	SeqID 1351	SeqID 1352	SeqID 1353

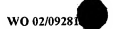
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1			į	Identities = 314/315 (99%), Positives = 315/315 (99%)
SeqID 1354	SA-2345.1	Contig135 (8014-8961 p)	96	[Streptococcus agalactiae] dbj[BAA82284.1] (AB028896) CpslaJ [Streptococcus agalactiae]
SeqID 1355	SA-2346.2	Contig135 (9045-10001 p)	86	Identities = 318/318 (100%), Positives = 318/318 (100%) dbj BAA33751.1  (AB017355) cpsJ [Streptococcus agalactiae] dbj BAA82285.1  (AB028896) CpsIaK [Streptococcus agalactiae] Length = 318
SeqID 1356	SA-2347.1	Contig75 (158-718 p)	51	Identities = 64/234 (27%), Positives = 122/234 (51%), Gaps = 3/234 (1%) spl032095 YUEF_BACSU HYPOTHETICAL 40.9 KDA PROTEIN IN DEGQ-ALD INTERGENIC REGION pir  G70007 conserved hypothetical protein yueF - Bacillus subtilis emb CAB15168.1  (299120) similar to hypothetical proteins [Bacillus subtilis] Length = 369
SeqID 1357	SA-2348.1	Contig75 (814-1494 p)	57	Identities = 90/210 (42%), Positives = 136/210 (63%) splQ02170 RADC_BACSU DNA REPAIR PROTEIN RADC HOMOLOG (ORFB) pir  B45239 DNA repair protein homolog ysxA - Bacillus subtilis gb AAA22396.1  (M96343) homologous to E. coli radC gene product and to unidentified protein from Staphylococcus aureus [Bacillus subtilis] gb AAA22583.1  (L08793) putative [Bacillus subtilis] emb CAB14764.1  (Z99118) similar to DNA repair protein [Bacillus subtilis] Length = 231
SeqID 1358	SA-2349.1	Contig75 (1507-2145 m)	49	Identities = 96/202 (47%), Positives = 140/202 (68%), Gaps = 4/202 (1%) pir  A69787 hypothetical protein ydiH - Bacillus subtilis dbj BAA19721.1  (D88802) ydiH [Bacillus subtilis] emb CAB12416.1  (Z99107) ydiH [Bacillus subtilis] Length = 215
SeqID 1359	SA-235.1	Contig136 (32600-32830 m)	No Hits found	
Sedin Ison	SA-2350.1	Contig/5 (2300-264/ m)	No Hits found	



Identities = 173/373 (46%), Positives = 236/373 (62%), Gaps = 6/373 (1%) dbj BAB04979.1  (AP001511) Fe-S cluster formation protein [Bacillus halodurans] Length = 386	Identities = 42/89 (47%), Positives = 63/89 (70%), Gaps = 2/89 (2%) gb AAF15359.1 AF201954_1 (AF201954)     phosphoribosylpyrophosphate synthetase [Plasmodium falciparum]   Length = 323	Identities = 48/70 (68%); Positives = 58/70 (82%) pir  T00087 rhamnosyltransferase - Streptococcus mutans db  BAA32090.1  (AB010970) rhamnosyltransferase [Streptococcus mutans] Length = 311	Identities = 234/362 (64%), Positives = 284/362 (77%) pir  T00086 rgpAc protein - Streptococcus mutans dbj BAA32089.1  (AB010970) rgpAc [Streptococcus mutans] Length = 362	Identitles = 257/283 (90%), Positives = 273/283 (95%) gb AAC38675.1  (AF030359) dTDP-L-rhamnose synthase (Streptococcus pneumoniae) gb AAC38685.1  (AF030361) dTDP-L-rhamnose synthase (Streptococcus pneumoniae) gb AAC38701.1  (AF030364) dTDP-L-rhamnose synthase (Streptococcus pneumoniae) gb AAD10184.1  (AF026471) Cps2O (Streptococcus pneumoniae)	Identities = 92/108 (85%), Positives = 100/108 (92%)   Abj BAA21508.1   (AB000631) unnamed protein product   Streptococcus mutans   Length = 111	Identities = 345/367 (94%), Positives = 358/367 (97%) dbj BAA21507.1  (AB000631) sigma 42 protein [Streptococcus mutans] Length = 371	Identities = 271/637 (42%), Positives = 389/637 (60%), Gaps = 56/637 (8%) splQ04505 PRIM_LACLA DNA PRIMASE pir  JC2485 DNA primase (EC 2.7.7) dnaG - Lactococcus lactis prf  2106154A DNA primase [Lactococcus lactis] Length =
   Identities = 173/373 (46%), Positive   6/373 (1%) dbj BAB04979.1  (AP001   protein [Bacillus halodurans]	.Identities = 42/89 (47%), Postal (2%) galAAF15359 phosphoribosylpyrophostal falciparum]	Identities = 48/70 (68%); rhamnosyltransferase - Str (AB010970) rhamnosyltl	Identities = 234/362 (t   pir  T00086 rgpAc p   dbj BAA32089.1  (AB010	Identities = 257/283 (90%), Positive gb AAC38675.1  (AF030359) dTDP-L [Streptococcus pneumoniae] gb AAC386 L-rhamnose synthase [Streptococ gb AAC38701.1  (AF030364) dTDP-L [Streptococcus pneumoniae] gb AAD Cps2O [Streptococcus pneumoniae]	identities = 92/108 (85%), Podb] BAA21508.1  (AB000631) [Streptococcus mutans]	Identities = 345/367 (9   dbj BAA21507.1  (AB0006   mutans	Identities = 271/637 (42%), Positives = 389/637 56/637 (8%) splQ04505 PRIM_LACLA DNA pir  JC2485 DNA primase (EC 2.7.7) dnaG - La pri  2106154A DNA primase [Lactococcus lactis]
63	89	82	99	. 85	82	83	64
··· Contig75 (2649-3767 m)	 Contig75 (3768-4043 m)	Contig72 (5600-5809 p)	Contig72 (4453-5610 p)	Contig72 (3485-4339 p)	Contig72 (3054-3395 p)	Contig72 (1803-2945 p)	Contig72 (20-1828 p)
SA-2351.1	SA-2352.1	SA-2353.1	SA-2354.1	SA-2355.1	SA-2356.1	SA-2357.1	SA-2358.1
SeqID 1361	SeqID 1362	SeqID 1363	SeqID 1364	SeqiD 1365	SeqID 1366	SeqID 1367	SeqID 1368

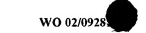


SeqID 1369	SA-2359.2	Contig71 (2968-3543 m)	93	identities = 96/195 (49%;, Positives = 138/195 (70%), Gaps = 15/195 (7%) gblAAK04732.1JAE006296_6 (AE006296) DNAdirected RNA polymerase delta chain (EC 2.7.7.6) [Lactococcus lactis subsp. lactis] Length = 187
SeqID 1370	SA-236.1	Contig136 (32214-32603 m)	No Hits found	
SeqID 1371	SA-2360.1	Contig71 (1091-2695 m)	68	Identities = 421/535 (78%), Positives = 481/535 (89%) emb CAA09021.2  (AJ010153) CTP synthetase [Lactococcus lactis subsp. cremoris] Length = 535
SeqID 1372	SA-2361.2	Contig71 (56-982 m)	. 29	Identities = 157/312 (50%), Positives = 212/312 (67%), Gaps = 9/312 (2%) gb AAK04219.1 AE006250_6 (AE006250) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 311
SeqID 1373	SA-2363.1	Contig104 (71-529 m)	No Hits found	
SeqID 1374	SA-2364.1	Contig104 (3-842 p)	88	Identities = 62/235 (26%), Positives = 108/235 (45%), Gaps = 12/235 (5%) pir  T34651 probable transmembrane protein - Streptomyces coelicolor emb CAA22372.1  (AL034446) putative transmembrane protein [Streptomyces coelicolor A3(2)] Length = 307
SeqID 1375	SA-2365.1	Contig104 (949-1539 p)	No Hits found	
SeqiD 1376	SA-2366.1	Contig104 (1573-2844 p)	47	Identities = 106/440 (24%), Positives = 206/440 (46%), Gaps = 65/440 (14%) pir[JC82901 conserved hypothetical UU367 [imported] - Ureaplasma urealyticum gb AAF30776 1 AE002133_9 (AE002133) conserved hypothetical [Ureaplasma urealyticum] Length = 507
SeqID 1377	SA-2367.1	Contig104 (2857-3288 p)	45	Identities = 39/145 (26%), Positives = 68/145 (46%), Gaps = 7/145 (4%) pir  D69831 conserved hypothetical protein yhfO - Bacillus subtilis emb CAA74538.1  (Y14084) hypothetical protein [Bacillus subtilis] emb CAB12871.1  (Z99109) similar to hypothetical proteins [Bacillus subtilis] Length = 149



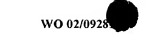
SeqID 1378	-SA-2368.1	Contig104 (3380-4264 p)	64	Identities = 145/283 (51%), Positives = 191/283 (67%), Gaps = 12/283 (4%) dbj BAB06129.1  (AP001515) tRNA pseudourldine 5S synthase [Bacillus halodurans] Length = 304
SeqID 1379	SA-2369.2	Contig104 (4277-5209 p)	86	Identities = 310/311 (99%), Positives = 311/311 (99%) gb AAB64408.1  (U92073) macrolide-efflux protein [Streptococcus agalactiae] Length = 311
SeqID 1380	SA-237.1	Contig136 (31877-32062 m)	No Hits found	
SeqID 1381	SA-2370.3	Contig124 (2287-4089 m)	59	Identities = 252/598 (42%), Positives = 358/598 (59%), Gaps = 54/598 (9%) gbjAAK04733.1jAE006296_7 (AE006296) conserved hypothetical protein [Lactococcus lactis subsp. lactis] Length = 550
SeqID 1382	SA-2371.1	Contig124 (1732-2214 m)	89	Identities = 86/154 (55%), Positives = 114/154 (73%), Gaps = 1/154 (0%) splP80240 GREA_BACSU TRANSCRIPTION ELONGATION FACTOR GREA (TRANSCRIPT CLEAVAGE FACTOR GREA) (GENERAL STRESS PROTEIN 20M) (GSP20M) pir JA69637 transcription elongation factor greA - Bacillus subtilis emb CAB14674.1  (299117) transcription elongation factor [Bacillus subtilis]
SeqID 1383	SA-2372.1	Contig124 (168-1628 m)	54	Identities = 182/488 (37%), Positives = 267/488 (54%), Gaps = 17/488 (3%) pirIIC75543 6-aminohexanoate-cyclic-dimer hydrolase - Deinococcus radiodurans (strain R1) gblAAF09821.1[AE001885_5 (AE001885) 6-aminohexanoate-cyclic-dimer hydrolase [Deinococcus radiodurans] Length = 561
SeqID 1384	SA-2373.1	Contig93 (16319-16855 p)	54	Identities = 65/206 (31%), Positives = 110/206 (52%), Gaps = 16/206 (7%) gb AA27630.1 (M98350) hydrophobic protein [unidentified bacterium]

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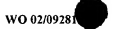
Identities = 54/192 (28%), Positives = 89/192 (46%), Gaps = 14/192 (7%) splP50736 YPDA_BACSU HYPOTHETICAL 36.3 KD PROTEIN IN RECQ-CMK INTERGENIC REGION pirl/A69934 thioredoxin reductase homolog ypdA - Bacillus subtilis gb AAC83954.1  (L47648) putative [Bacillus subtilis] emb CAB14227.1  (Z99116) similar to thioredoxin reductase [Bacillus subtilis] Length = 324	Identities = 213/322 (66%), Positives = 260/322 (80%), Gaps = 5/322 (1%) gb AAB81912.1  (U92974) unknown [Lactococcus lactis]		Identities = 174/321 (54%), Positives = 241/321 (74%), Gaps = 4/321 (1%) pir  JC5050 sugar phosphate transport protein - Shigella flexneri gb AAC44575.1  (U28354) IS629 ORFB fused with sequences similar to E. coli GlpT and UhpT proteins, Swiss-Prot Accession Number P08194 and P09836; Method: conceptual translation supplied by author [Shigella flexneri] Length = 333	Identities = 23/57 (40%), Positives = 36/57 (62%) ref[NP_049417.1] putative cro-like regulatory protein [Streptococcus thermophilus bacteriophage DT1] gb[AAD21905.1] (AF085222) putative cro-like regulatory protein [Streptococcus thermophilus bacteriophage DT1] Length = 67	Identities = 112/253 (44%), Positives = 161/253 (63%), Gaps = 1/253 (63%), Gaps = 1/253 (0%) splQ9RGS6JTHIM_STACA HYDROXYETHYLTHIAZOLE KINASE (4-METHYL-5-BETA-HYDROXYETHYLTHIAZOLE KINASE) (THZ KINASE) (THZ KINASE) (TH KINASE) gbjAAF25543.1 AF109218_3 (AF109218) Thim [Staphylococcus carnosus]
2	7	No Hits found	υ 		, v
Contig93 (15061-16188 m)	Contig93 (14038-15024 p)	Contig93 (13570-13875 m)	Contig93 (12039-13412 p)	Contig136 (31331-31807 m)	Contig130 (2285-3055 p)
SA-2374.1	SA-2376.1	SA-2377.1	SA-2378.2	SA-238.1	SA-2380.2
SeqID 1385	SeqID 1386	SeqID 1387	SeqID 1388	SeqID 1389	SeqID 1390

SeqID :1391	. SA-2381.1	Contig130 (1486-2283 p)	67	Identities = 139/258 (53%), Positives = 186/258 (71%), Gaps = 4/258 (1%) gb AAF25542.1 AF109218 <u>-2</u> (AF109218) ThiD [Staphylococcus carnosus]
SeqID 1392	SA-2382.1	Contig130 (805-1461 p)	52	Identities = 74/213 (34%), Positives = 122/213 (56%), Gaps = 13/213 (6%) gb AAF25541.1 AF109218_1 (AF109218) TenA [Staphylococcus carnosus] Length = 228
SeqID 1393	SA-2383.1	Contig130 (181-306 p)	No Hits found	
SeqID 1394	SA-2384.1	Contig130 (20-721 p)	4	Identities = 48/216 (22%), Positives = 98/216 (45%), Gaps = 3/216 (1%) emb CAA91230.1  (256283) orf2 [Lactobacillus helveticus] Length = 217
SeqID 1395	SA-2386.1	Contig101 (9386-10810 p)	64	Identities = 189/462 (40%), Positives = 313/462 (66%) emb[CAB52225.1  (298171) EpsU protein [Streptococcus thermophilus] Length = 471
SeqID 1396	SA-2387.1	Contig101 (8022-9386 p)	No Hits found	
SeqID 1397	SA-2388.2	Contig101 (7073-8020 p)	23	Identities = 53/116 (45%), Positives = 75/116 (63%), Gaps = 4/116 (3%) gb AAF18951.1 AF155805_5 (AF155805) Cps9H [Streptococcus suis]
SeqID 1398	SA-239.1	Contig136 (30561-31331 m)	88	Identities = 99/260 (38%), Positives = 157/260 (60%), Gaps = 3/260 (1%) pir  S45085 hypothetical protein zeta - Streptococcus pyogenes plasmid pDB101 and pBT233 pir  S68606 hypothetical protein zeta - Streptococcus pyogenes plasmid pBT233 emb CA45934.1  (X64695) ORF zeta [Streptococcus pyogenes] emb CA47091.1  (X66468) orf zeta [Streptococcus pyogenes] emb CA47092.1  (X66468) orf zeta [Streptococcus pyogenes] = 287

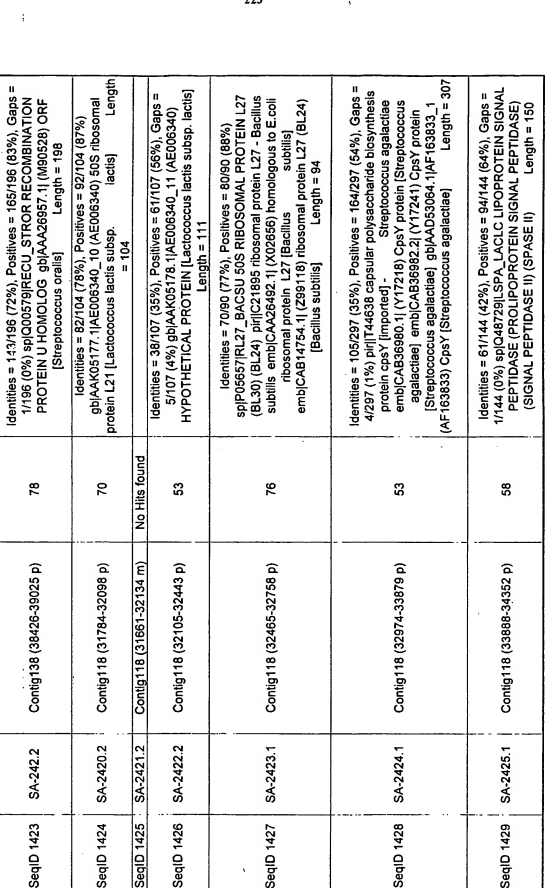


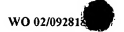
SeqID 1399	SA-2390.2	Contig136 (91497-93569 p)	38	Identities = 178/535 (33%), Positives = 269/535 (50%), Gaps = 55/535 (10%) spiP54602 YHCR_BACSU HYPOTHETICAL 132.7 KDA PROTEIN IN CSPB-GLPP INTERGENIC REGION pirt  F69823 probable phosphoesterase (EC 3.1) yhcR - Bacillus subtilis emb  CAB12747.1  (X96983) hypothetical protein [Bacillus subtilis] emb  CAB12747.1  (Z99108) similar to 5-nucleotidase [Bacillus subtilis]
SeqID 1400	SA-2391.1	Contig136 (93606-94016 m)	02	Identities = 72/136 (52%), Positives = 96/136 (69%) splo08450IDEF_CLOBE POLYPEPTIDE DEFORMYLASE (PDF) (FORMYLMETHIONINE DEFORMYLASE) emblCAB09662.11 (Z96934) peptide deformylase [Clostridium beijerinckii] Length = 136
SeqID 1401	SA-2392.1	Contig136 (94086-95393 m)	81	Identities = 292/436 (66%), Positives = 356/436 (80%), Gaps = 2/436 (0%) dbj BAB05820.1  (AP001514) NADP-specific glutamate dehydrogenase [Bacillus halodurans] Length = 458
SeqID 1402	SA-2394.2	Contig89 (9769-11001 m)	59	Identities = 63/243 (25%), Positives = 120/243 (48%) dbj BAB03800.1  (AP001507) BH0081~unknown conserved protein in others [Bacillus halodurans] Length = 251
SeqID 1403	SA-2395.1	Contig89 (11041-12582 m)	8	Identities = 352/509 (69%), Positives = 421/509 (82%), Gaps = 1/509 (0%) pir  D69813 ABC transporter (ATP-binding protein) homolog yfmM - Bacillus subtilis dbj BAA22327.1  (D86417) YfmM [Bacillus subtilis] emb CAB12571.1  (299108) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis] Length = 518
SeqID 1404	SA-2396.2	Contig88 (2596-3666 p)	09	Identities = 138/358 (38%), Positives = 220/358 (60%), Gaps = 18/358 (5%) pir E71373 probable regulatory protein (pfoS/R) - syphilis spirochete gb AAC65034.1  (AE001189) regulatory protein (pfoS/R) [Treponema pallidum] Length = 350

	Identities = 163/325 (50%), Positives = 222/325 (68%), Gaps = 3/325 (0%) dbj BAB07127.1] (AP001518) thioredoxin reductase [Bacillus halodurans] Length = 330	Identities = 144/246 (58%), Positives = 186/246 (75%), Gaps = 6/246 (2%) dbj BAB06198.1  (AP001515) tRNA methyltransferase	1/174 (50%), Positives = 128/174 (72%), Gaps = 1/174 (0%) spjO31740 RIMM_BACSU PROBABLE 16S RRNA PROCESSING PROTEIN RIMM pir E69880 conserved hypothetical protein ylqE - Bacillus subtilis emb CAB13475.1  (299112) similar to hypothetical proteins [Bacillus subtilis]	Identities = 65/236 (27%), Positives = 117/236 (49%), Gaps = 27/236 (11%) pir][H83403 hypothetical protein PA1939 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG05327.1 AE004620_6 (AE004620) hypothetical protein [Pseudomonas aeruginosa] Length = 665	1dentities = 412/632 (65%), Positives = 506/632 (79%), Gaps = 6/632 (0%) pir  S68599 phosphotransferase system enzyme II (EC 2.7.1.69), sucrose-specific - Streptococcus sobrinus (strain 6715) Length = 632	ldentities = 252/254 (99%), Positives = 253/254 (99%) 405.1 Contig70 (5129-5893 m) 98 gb AAG09975.1 AF248038_4 (AF248038) methyltransferase [Streptococcus agalactiae] Length = 254	Identities = 112/112 (100%), Positives = 112/112 (100%)   406.1   Contig70 (4770-5108 m)   95   gb AAG09974.1 AF248038_3 (AF248038) unknown (Streptococcus agalactiae)   Length = 112	ldentities = 101/101 (100%), Positives = 101/101 (100%) 407.1 Contig70 (4463-4768 m) 94 gb AAG09973.1 AF248038_2 (AF248038) acetate kinase
	SA-2397.1	SA-2398.1	SA-2399.1	SA-240.2	SA-2404.1	SA-2405.1	SA-2406.1	SA-2407.1



SeqID 1413	SA-2409.2	Contig70 (3706-4245 p)	52	Identities = 105/107 (98%), Positives = 106/107 (98%)   gb AAG09972.1 AF248038_1 (AF248038) integrase   Streptococcus agalactiae]
SeqID 1414	SA-241.2	Contig138 (37833-38351 m)	<b>.</b>	Identities = 74/175 (42%), Positives = 98/175 (55%), Gaps = 6/175 (3%) splP50838 YPSA_BACSU HYPOTHETICAL 21.1 KD PROTEIN IN COTD-KDUD INTERGENIC REGION pir  D69941 conserved hypothetical protein ypsA - Bacillus subtilis gb AAB38471.1  (L47838) putative [Bacillus subtilis] emb CAB14136.1  (299115) similar to hypothetical proteins from B. subtilis [Bacillus subtilis]
SeqID 1415	SA-2410.1	Contig70 (3110-3544 p)	42	Identities = 40/123 (32%), Positives = 63/123 (50%), Gaps = 9/123 (7%) gb AAD00268.1  (U77495) putative integrase [Leuconostoc cenos bacteriophage 10MC]
SeqID 1416	SA-2411.1	Contig70 (2734-3060 p)	33	Identities = 24/72 (33%), Positives = 38/72 (52%), Gaps = 2/72 (2%) dbj BAB07266.1  (AP001519) unknown conserved protein in others [Bacillus halodurans] Length = 79
SeqID 1417	SA-2412.1	Contig70 (2007-2561 p)	No Hits found	
SeqID 1418	SA-2414.1	Contig70 (1022-1783 p)	38	Identities = 70/211 (33%), Positives = 101/211 (47%), Gaps = 5/211 (2%) pir JH0204 hypothetical 30.5K protein precursor - Enterococcus faecalis plasmid pAM-beta-1 gb AAC38600.1  (AF007787) orfC [Enterococcus faecalis] Length = 288
SeqID 1419	SA-2415.1	Contig70 (440-997 p)	No Hits found	
SeqID 1420	SA-2416.1	Contig70 (12-440 p)	No Hits found	
SeqID 1421	SA-2418.2	Contig104 (16088-16768 p)	88	Identities = 202/246 (82%), Positives = 223/246 (90%), Gaps = 7/246 (2%) emb CAB54564.1  (AJ005926) response regulator [Streptococcus pneumoniae] Length = 246
SeqID 1422	SA-2419.2	Contig104 (13377-15926 p)	92	Identities = 555/847 (65%), Positives = 673/847 (78%), Gaps = 4/847 (0%) emb[CAB50785.1] (AJ007700) aminopeptidase N [Streptococcus thermophilus] Length = 847

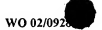




-SeqID 1430	SA-2426.2	Contig118 (34336-34836 p)	80	Identities = 115/168 (68%), Positives = 140/168 (82%)   gb AAK05096.1 AE006334_2 (AE006334) pseudouridine   synthase [Lactococcus lactis subsp.   lactis]   Length = 301
SeqID 1431	SA-2428.1	Contig67 (20-2098 p)	87	Identities = 536/692 (77%), Positives = 613/692 (88%) emb[CAC09927.1  (AJ249559) translation elongation factor G, EF G [Bacillus stearothermophilus] Length = 692
SeqID 1432	SA-2429.2	Contig135 (26612-27559 m)	23	Identities = 42/173 (24%), Positives = 80/173 (45%), Gaps = 2/173 (1%) refINP_052734.1  pXO1-38 [Bacillus anthracis] pir[ F59095 hypothetical protein pXO1-38 - Bacillus anthracis virulence plasmid pXO1 gb AAD32342.1 AAD32342 (AF065404) pXO1-38 [Bacillus anthracis] Length = 276
SeqID 1433	SA-2430.1	Contig135 (25417-26493 m)	64	Identitles = 153/350 (43%), Positives = 234/350 (66%), Gaps = 4/350 (1%) ref[NP_049990.1] orf359 gp [Streptococcus thermophilus bacteriophage Sfi21] emb[CAA64931.1] (X95646) integrase [Streptococcus thermophilus bacteriophage Sfi21] gb[AAC03454.1] (AF020798) integrase homolog [Streptococcus thermophilus bacteriophage TP-J34] gb[AAD44095.1]AF115103_25 (AF115103) orf359 gp [Streptococcus thermophilus bacteriophage Sfi21] Length = 359
SeqID 1434	SA-2431.3	Contig135 (23774-25237 p)	63	Identities = 251/401 (62%), Positives = 312/401 (77%), Gaps = 4/401 (0%) gblAAK04936.1 AE006317_5 (AE006317) 30S ribosomal protein S1 [Lactococcus lactis subsp. lactis] Length = 408

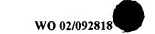


SeqID 1435	SA-2432.3	Contig135 (24256-25128 m)	38	Identities = 73/265 (27%), Positives = 111/265 (41%), Gaps = 40/265 (15%) refINP_01442.1  anchorage subunit of a-agglutinin; Aga1p [Saccharomyces cerevisiae] sp P32323 AGA1_YEAST A-AGGLUTININ ATTACHMENT SUBUNIT PRECURSOR pir  A41258 a-agglutinin core protein AGA1 - yeast (Saccharomyces cerevisiae) gb AAA34382.1  (M60590) a-agglutinin core subunit [Saccharomyces cerevisiae] emb CAA96325.1  (Z71659) ORF YNR044w [Saccharomyces cerevisiae]
SeqiD 1436	SA-2433.2	Contig96 (1706-2137 m)	42	Identities = 47/137 (34%), Positives = 71/137 (51%), Gaps = 5/137 (3%) dbjjBAB04953.1  (AP001511) small multidrug export related protein [Bacillus halodurans] Length = 134
SeqID 1437	SA-2434.3	Contig96 (507-1547 p)	81	Identities = 242/338 (71%), Positives = 290/338 (85%)   gb AAK06250.1 AE006444_1 (AE006444) elongation factor Ts   Lactococcus lactis subsp. lactis
SeqID 1438	SA-2435.3	Contig96 (3-413 p)	83	Identities = 102/131 (77%), Positives = 115/131 (86%), Gaps = 2/131 (1%) gblAAK06251.1 AE006444_2 (AE006444) 30S ribosomal protein S2 [Lactococcus lactis subsp. lactis]
SeqID 1439	SA-2436.3	Contig96 (2-316 m)	30	Identities = 28/59 (47%), Positives = 32/59 (53%) emb[CAB67155.1  (AJ271079) hypothetical protein [Oenothera elata subsp. hookeri]
SeqiD 1440	SA-2437.2	Contig135 (2125-2817 p)	6	Identities = 229/230 (99%), Positives = 229/230 (99%) splQ04662lCPSB_STRAG CPSB PROTEIN pir  S34975 polysaccharide chain length regulator cpsB [imported] - Streptococcus agalactiae gb AAB00362.1  (AF163833) CpsC [Streptococcus agalactiae] Length = 230
SeqID 1441	SA-2438.1	Contig135 (1385-2116 p)	96	Identities = 242/243 (99%), Positives = 243/243 (99%)   dbj BAA82276.1  (AB028896) CpslaB [Streptococcus agalactiae]   Length = 243
SeqID 1442	SA-2439.1	Contig135 (80-427 m)	No Hits found	



SeqID 1443	SA-244.2	Contig138 (39012-41258 p)	71	Identities = 412/725 (56%), Positives = 537/725 (73%), Gaps = 25/725 (3%) gb AAF17262.1 AF210752_1 (AF210752) penicillinbinding protein 1A [Streptococcus pneumoniae]
SeqID 1444	SA-2440.1	Contig135 (12-1379 p)	66	Identities = 450/453 (99%), Positives = 452/453 (99%) emb CAB36981.1  (Y17218) CpsX protein [Streptococcus agalactiae] dbj BAA82275.1  (AB028896) CpsIaA [Streptococcus agalactiae] Length = 485
SeqID 1445	SA-2441.1	Contig123 (43050-43538 m)	No Hits found	
SeqID 1446	SA-2442.1	Contig123 (42359-42814 p)	No Hits found	
SeqID 1447	SA-2443.1	Contig123 (41386-42402 p)	No Hits found	
SeqiD 1448	SA-2445.2	Contig123 (39711-40976 p)	84	Identities = 138/398 (34%), Positives = 208/398 (51%), Gaps = 17/398 (4%) refINP_076752.1 integrase [bacteriophage blL310] gb AAK04145.1 AE006243_9 (AE006243) prophage ps1 protein. 23, integrase [Lactococcus lactis subsp. lactis] gb AAK08405.1 AF323671_1 (AF323671) integrase [bacteriophage blL310] Length = 394
SeqID 1449	SA-2446.2	Contig136 (2790-3302 m)	48	Identities = 78/191 (40%), Positives = 112/191 (57%), Gaps = 9/191 (4%) dbj BAA21095.1 (D88438) repB [Lactobacillus acidophilus] Length = 193
SeqID 1450	SA-2447.1	Contig136 (2553-2756 m)	No Hits found	
SeqID 1451	SA-2448.1	Contig136 (1354-2535 m)	4	Identities = 101/370 (27%), Positives = 167/370 (44%), Gaps = 39/370 (10%) pir  T13283 probable integrase - Streptococcus phage phi-O1205 gb AAC79517.1  (U88974) ORF1 [Streptococcus thermophilus temperate bacteriophage O1205] Length = 359
SeqID 1452	SA-2450.2	Contig135 (78598-79290 m)	No Hits found	
SeqID 1453	SA-2452.1	Contig66 (2566-2703 m)	No Hits found	

Identities = 112/244 (45%), Positives = 164/244 (66%), Gaps = 10/244 (4%) spl031458 YBFT_BACSU HYPOTHETICAL 27.3 KD PROTEIN IN GLTP-CWLJ INTERGENIC REGION pir  E69750 glucosamine-6-phosphate isomerase homolog ybfT - Bacillus subtilis emb CAB12030.1  (299105) similar to glucosamine-6-phosphate isomerase [Bacillus subtilis]	liis] /318 (7 sp. (sti	8 (75% (strain) 1 protei 3 (wouric udouric = 238 3%), Ge	Identities = 33/83 (39%), Positives = 51/83 (60%) pir  E83144 hypothetical protein PA4016 [imported] - Pseudomonas
Identities = 112/2 10/244 (4%) spl( KD PROTEII pir  E69750 glucos Bacillus subt	dbj BAA33133.1  ( Identities = 1 pir  S76960 hypot 6803) dbj B,	Identities = 1  pir  S76960 hypot 6803) dbj  B. [Syne Identities = 91/19 Synthase [E 57/620 (9%))	Identities = 33/83 (39%), Positives = 51/83 (60%) pir  E83144 hypothetical protein PA4016 [imported] - Pseudomonas (strain PAO1) gb AAG07403.1 AE004818
. 67	70	50 60	39
Contig66 (1714-2415 m)	Contig66 (686-1642 p)	Contig66 (686-1642 p)  Contig66 (3-590 m)  Contig132 (33811-35607 p)	Contig132 (33590-33832 p)
SA-2453.1	SA-2454.1	SA-2454.1 SA-2455.1 SA-2457.3	SA-2458.3
SeqID 1454	SeqID 1455	SeqID 1455 SeqID 1456 SeqID 1457	SeqID 1458



SeqID 1460	SA-246.1	Contig138 (41304-42638 m)	06	Identities = 363/445 (81%), Positives = 408/445 (91%) splQ56115 PEPC_STRTR AMINOPEPTIDASE C pir  S48143 cysteine aminopeptidase C - Streptococcus thermophilus emb CAA82960.1  (Z30315) aminopeptidase C [Streptococcus thermophilus]
SeqID 1461	SA-2460.1	Contig91 (4551-5126 p)	56	Identities = 97/188 (51%), Positives = 133/188 (70%) gb AAC18360.1  (AF064763) putative membrane spanning protein [Lactococcus lactis subsp. cremoris] Length = 196
SeqID 1462	SA-2461.1	Contig91 (3911-4414 p)	No Hits found	
SeqID 1463	SA-2462.1	Contig91 (3130-3873 p)	59	Identities = 106/246 (43%), Positives = 150/246 (60%), Gaps = 7/246 (2%) pir  G69984 rRNA methylase homolog ysgA - Bacillus subtilis emb CAA99602.1  (Z75208) hypothetical protein [Bacillus subtilis] emb CAB14825.1  (Z99118) similar to rRNA methylase [Bacillus subtilis] Length = 248
SeqID 1464	SA-2463.1	Contig91 (2820-3377 m)	29	Identities = 35/91 (38%), Positives = 54/91 (58%), Gaps = 3/91 (3%) pir] G72240 hypothetical protein TM1564 - Thermotoga maritima (strain MSB8) gb AAD36630.1 AE001801_17 (AE001801) acylphosphatase, putative [Thermotoga maritima] Length = 90
SeqID 1465	SA-2464.2	Contig91 (1803-2693 m)	63	Identities = 140/307 (45%), Positives = 201/307 (64%), Gaps = 19/307 (6%) gb AAK04667.1 AE006291_1 (AE006291) conserved hypothetical protein [Lactococcus lactis subsp.   lactis] Length = 307

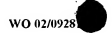
SeqiD 1466	SA-2465.1	- Contig139 (189466-189870 m)	54	Identities = 42/105 (40%), Positives = 62/105 (59%) sp[P45678 PEB1_CAMJE MAJOR CELL-BINDING FACTOR PRECURSOR (CBF1) (PEB1) pir  A48518 probable ABC-type amino-acid transporter periplasmic solute-binding protein (50921c precursor [imported] - Campylobacter jejuni (strain NCTC 11168) gb[AAA02919.1] (L13662) major cell-binding factor [Campylobacter jejuni] emb[CAB73178.1] (AL139076) probable ABC-type amino-acid transporter periplasmic solute-binding protein [Campylobacter jejuni]
SeqID 1467	SA-2466.1	Contig139 (188758-189453 m)	89	Identities = 112/226 (49%), Positives = 161/226 (70%), Gaps = 3/226 (1%) pir  G81365 probable ABC-type amino-acid transporter permease protein Cj0920c [imported] - Campylobacter jejuni (strain NCTC 11168) emb CAB73177.1  (AL139076) putative ABC-type amino-acid transporter permease protein [Campylobacter jejuni] Length = 250
SeqID 1468	SA-2467.1	Contig139 (188096-188746 m)	. 88	Identities = 85/216 (39%), Positives = 132/216 (60%), Gaps = 6/216 (2%) pir  F69633 glutamine ABC transporter (membrane protein) glnP - Bacillus subtilis emb CAB14687.1  (Z99117) glutamine ABC transporter (integral membrane protein) [Bacillus subtilis] emb CAB14704.1  (Z99118) glutamine ABC transporter (integral membrane protein) [Bacillus subtilis]
SeqID 1469	SA-2468.2	Contig139 (187481-188047 p)	42	Identities = 43/157 (27%), Positives = 83/157 (52%), Gaps = 9/157 (5%) dbj BAB04094.1  (AP001508) BH0375~unknown conserved protein in B. subtilis [Bacilius halodurans] Length = 194
SeqID 1470	SA-2469.2	Contig139 (186295-187314 p)	No Hits found	



SeqID 1471	SA-247.1	Contig138 (42751-43572 m)		Identities = 173/275 (62%), Positives = 215/275 (77%), Gaps = 1/275 (0%) splP18843 NADE_ECOLI NH(3)-DEPENDENT NAD(+) SYNTHETASE (NITROGEN-REGULATORY PROTEIN) pir  D64933 NAD+ synthase (EC 6.3.1.5) nadE [validated] - Escherichia coli (strain K-12) dbj  BAA15529.1  (D90817) NH(3)-dependent NAD(+) synthetase (EC 6.3.5.1) (Nitrogen-regulatory protein). [Escherichia coli] dbj  BAA15535.1  (D90818) NH(3)-dependent NAD(+) synthetase (EC 6.3.5.1) (Nitrogen-regulatory protein). [Escherichia coli] gb  AAC74810.1  (AE000269) NAD synthetase, prefers NH3 over glutamine [Escherichia coli K12]
SeqID 1472	SA-2470.1	Contig65 (2150-2422 m)	No Hits found	
SeqID 1473	SA-2471.1	Contig65 (1678-2109 p)	67	Identities = 69/152 (45%), Positives = 101/152 (66%), Gaps = 12/152 (7%) pir[ T11571 hypothetical protein 2 - Streptococcus mutans gb AAD15622.1  (U75480) unknown [Streptococcus mutans] Length = 151
SeqID 1474	SA-2472.1	Contig65 (1283-1681 p)	77	Identities = 88/129 (68%), Positives = 112/129 (86%) pir  T11570 hypothetical protein 1 - Streptococcus mutans gb AAD15621.1  (U75480) unknown [Streptococcus mutans] Length = 131
SeqID 1475	SA-2473.1	Contig65 (495-1268 p)	85	Identities = 184/258 (71%), Positives = 227/258 (87%) splP72482 LGT_STRMU PROLIPOPROTEIN DIACYLGLYCERYL TRANSFERASE pir  T11569 prolipoprotein diacylglyceryl transferase (EC 2.4.99) Streptococcus mutans gb AAC80171.3  (U75480) putative prolipoprotein diacylglycerol transferase [Streptococcus mutans] Length = 259
SeqID 1476	SA-2474.1	Contig65 (377-502 p)	88	Identities = 68/77 (88%), Positives = 75/77 (97%) sp Q9ZA98 HPRK_STRSL HPR(SER) KINASE/PHOSPHATASE gb AAD12781.1  (AF069743) HPr(serine) kinase [Streptococcus salivarius] Length = 309



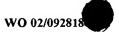
SeqID 1477	SA-2475.1	Contig65 (1-264 p)	95	Identities = 82/84 (97%), Positives = 84/84 (99%) dbj BAA77782.1  (AB027460) Hpr kinase [Streptococcus bovis] Length = 310
SeqID 1478	SA-2476.1	Contig64 (2858-3013 m)	75	Identities = 33/46 (71%), Positives = 39/46 (84%) pir  H70958 probable uracil phosphoribosyltransferase (EC 2.4.2.9) - Mycobacterium tuberculosis (strain H37RV) emb CAB02640.1  (Z81011) pyrR [Mycobacterium tuberculosis]
SeqID 1479	SA-2477.1	Contig64 (1739-2809 m)	72	Identities = 188/352 (53%), Positives = 265/352 (74%) emb CAB89872.1  (AJ132624) carbamoyl phosphate synthetase small subunit [Lactococcus lactis] Length = 357
SeqID 1480	SA-2479.1	Contig64 (85-1683 m)	37	Identities = 116/414 (28%), Positives = 204/414 (49%), Gaps = 31/414 (7%) spiP77886 CARB_LACPL CARBAMOYL-PHOSPHATE SYNTHASE, PYRIMIDINE-SPECIFIC, LARGE CHAIN (CARBAMOYL-PHOSPHATE SYNTHETASE AMMONIA CHAIN) emb CAA91005.1  (Z54240) carbamoyl-phosphate synthase [Lactobacillus plantarum] Length = 1058
SeqID 1481	SA-248.1	Contig138 (43569-45029 m)	77	Identities = 309/476 (64%), Positives = 384/476 (79%), Gaps = 2/476 (0%) pir  D70008 nicotinate phosphoribosyltransferase homolog yueK - Bacillus subtilis emb CAB15163.1  (Z99120) similar to nicotinate phosphoribosyltransferase [Bacillus subtilis] Length = 490
SeqID 1482	SA-2480.2	Contig79 (5026-5787 p)	09	Identities = 125/247 (50%), Positives = 187/247 (75%) pir  D69763 ferrichrome ABC transporter (ATP-binding p) homolog yclP - Bacillus subtilis dbj BAA09014.1  (D50453) homologue of iron dicitrate transport ATP-binding protein FecE of E. coli [Bacillus subtilis] emb CAB12190.1  (Z99106) similar to ferrichrome ABC transporter (ATP-binding protein) [Bacillus subtilis] Length = 252



1	
Contig138 (45187-46101 m)  Contig69 (3312-4022 m)  Contig69 (2050-3297 m)	SA-248.2 Contig69 (4025-5050 SA-249.1 Contig138 (45187-4610 SA-2490.2 Contig69 (3312-4022 SA-2491.2 Contig69 (2050-3297)
	SA-2488.2 SA-2490.2 SA-2491.2

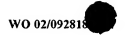


SeqID 1490	SA-2492.2	Contig106 (19268-21082 p)	. 22	Identities = 356/604 (58%), Positives = 445/604 (72%), Gaps = 4/604 (0%) sp[P39754 GLMS_BACSU GLUCOSAMINE—FRUCTOSE-6-PHOSPHATE AMINOTRANSFERASE [ISOMERIZING] (HEXOSEPHOSPHATE AMINOTRANSFERASE) (GFAT) (L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE AMIDOTRANSFERASE) (GLUCOSAMINE-6-PHOSPHATE AMIDOTRANSFERASE) (GLUCOSAMINE-6-PHOSPHATE SYNTHASE) pirilB69633 glutamine-fructose-6-phosphate transaminase (isomerizing) (EC 2.6.1.16) glmS - Bacillus subtilis gb[AAA64224.1] (U21932) L-glutamine-D-fructose-6-phosphate amidotransferase [Bacillus subtilis] emb[CAB11954.1] (Z99104) L-glutamine-D-fructose-6-phosphate amidotransferase [Bacillus subtilis] db][BAA33071.1] (AB006424) L-GLUTAMINE-D-FRUCTOSE-6-PHOSPHATE AMIDO TRANSFERASE [Bacillus subtilis] Length = 600
SeqID 1491	SA-2493.1	Contig62 (1978-3021 p)	83	Identities = 255/348 (73%), Positives = 298/348 (85%), Gaps = 1/348 (0%) sp[P10539[DHAS_STRMU ASPARTATE-SEMIALDEHYDE DEHYDROGENASE (ASADEHYDROGENASE) (ASADH) pir[]A29137 aspartate-semialdehyde dehydrogenase (EC 1.2.1.11) - Streptococcus mutans gb[AAA26850.1] (J02667) aspartate beta-semialdehyde dehydrogenase (EC 1.2.1.11) [Streptococcus mutans]
SeqID 1492	SA-2494.1	Contig62 (640-1494 m)	34	Identities = 71/271 (26%), Positives = 97/271 (35%), Gaps = 5/271 (1%) pirj S54157 extensin-like protein - cowpea (fragment)
SeqID 1493	SA-2495.1	Contig62 (537-1769 p)	42	Identities = 86/373 (23%), Positives = 176/373 (47%), Gaps = 26/373 (6%) gb AAF48863.1  (AE003509) CG15040 gene product [Drosophila melanogaster] Length = 1895
SeqID 1494	SA-2497.2	Contig104 (5252-5665 p)	99	Identities = 74/126 (58%), Positives = 101/126 (79%)   gb AAF21893.1 AF103794_1 (AF103794) unknown [Listeria   monocytogenes]



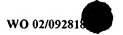
SeqID 1495	SA-2498.2	Contig104 (5658-5939 p)	47	Identities = 33/78 (42%), Positives = 50/78 (63%) pir  C69864 hypothetical protein yktA - Bacillus subtilis emb CAB13337.1  (Z99111) yktA [Bacillus subtilis] gb AAC24938.1  (AF012285) unknown [Bacillus subtilis]
SeqiD 1496	SA-2499.3	Contig104 (5929-6693 p)	56	Identities = 121/252 (48%), Positives = 171/252 (67%), Gaps = 4/252 (1%) gb AAK04646.1 AE006288_9 (AE006288) conserved hypothetical protein [Lactococcus lactis subsp. lactis] Length = 255
SeqID 1497	SA-25.1	Contig137 (15915-20645 p)	#	Identities = 117/377 (31%), Positives = 182/377 (48%), Gaps = 73/377 (19%) ref[NP_053235.1  pXO2-81 [Bacillus anthracis] gb[AAF13685.1 AF188935_83 (AF188935) pXO2-81 [Bacillus anthracis] Length = 589
SeqID 1498	SA-250.1	Contig138 (46170-46418 m)	No Hits found	
SeqID 1499	SA-2500.3	Contig104 (6761-8071 p)	48	Identities = 139/466 (29%), Positives = 213/466 (44%), Gaps = 36/466 (7%) pir  A82193 Sun/nucleolar protein family protein VC1502 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF94657.1  (AE004228) Sun/nucleolar protein family protein [Vibrio cholerae] Length = 503
SeqID 1500	SA-2501.2	Contig63 (2458-3795 m)	47	Identities = 111/428 (25%), Positives = 210/428 (48%), Gaps = 39/428 (9%) pir  B69855 amino acid permease homolog ykbA - Bacillus subtilis emb CAA05566.1  (AJ002571) YkbA [Bacillus subtilis] emb CAB13143.1  (Z99110) similar to amino acid permease [Bacillus subtilis] Length = 438
SeqID 1501	SA-2502.2	Contig63 (1737-2285 m)	17	Identities = 111/168 (66%), Positives = 136/168 (80%), Gaps = 3/168 (1%) gb AAK05247.1 AE006347_5 (AE006347) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 169

Identities = 130/381 (34%), Positives = 221/381 (57%), Gaps = 4/381 (1%) sp:Q08432 PATB_BACSU PUTATIVE   AMINOTRANSFERASE B pir[ S32934 aminotransferase patB - Bacillus subtilis   emb[CAB07910.1] (293933) aminotransferase   [Bacillus subtilis] emb[CAB07924.1] (293934) aminotransferase   [Bacillus subtilis] emb[CAB15133.1] (299120) aminotransferase   Bacillus subtilis] emb[CAB15133.1] (299120) aminotransferase   Bacillus subtilis]   Length = 387	9-322 m) 84 gb AAA71926.1  (L08445) uracil phosphoribosyltransferase [Streptococcus salivarius] Length = 128	Identities = 49/60 (81%), Positives = 54/60 (89%) 90 gb AAA71926.1  (L08445) uracil phosphoribosyltransferase [Streptococcus salivarius] Length = 128	Identities = 27/75 (36%), Positives = 52/75 (69%) refiNP_050766.1j acyl carrier protein [Guillardia theta] splP29189 ACP_GUITH ACYL CARRIER PROTEIN gb AAC35700.1  (AF041468) acyl carrier protein [Guillardia theta] Length = 81	Identities = 174/330 (52%), Positives = 239/330 (71%), Gaps = 2/330 (0%) splP71018 PLSX_BACSU FATTY ACID/PHOSPHOLIPID SYNTHESIS PROTEIN PLSX ACID/PHOSPHOLIPID SYNTHESIS PROTEIN PLSX pir  H69679 involved in fatty acid/phospholipid synthesis plsX - Bacillus subtilis emb CAA74248.1  (Y13937) putative PlsX protein [Bacillus subtilis] emb CAB13462.1  (299112) attemate
Contig63 (432-1601 m)	Contig63 (179-322 m)	Contig63 (3-182 m)	Contig132 (1877-2116 p)	Contig132 (874-1866 p)
SA-2503.1	SA-2503.1 SA-2504.1		SA-2507.2	SA-2508.1
SeqID 1502	SeqID 1503	SeqID 1504	SeqID 1505	SeqiD 1506



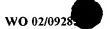
=	E		Φ .	
Identities = 121/247 (48%), Positives = 176/247 (70%) spl034900 YTMN_BACSU PROBABLE AMINO-ACID ABC TRANSPORTER ATP-BINDING PROTEIN YTMN pir  F69641 histidine transport protein hisP - Bacillus subtilis gb AAC00329.1 (AF008220) putative amino acid transporter [Bacillus subtilis] emb CAB14894.1 (Z99118) histidine transport protein (ATP-binding protein) [Bacillus subtilis] emb CAB14912.1 (Z99119) histidine transport protein (ATP-binding protein)	Identities = 430/659 (65%), Positives = 537/659 (81%), Gaps = 4/659 (0%) gb AAK04722.1 AE006295_3 (AE006295) potassium uptake protein [Lactococcus lactis subsp. lactis]  Length = 671	Identities = 142/248 (57%), Positives = 179/248 (71%), Gaps = 1/248 (0%) spIP45200 YDFG_HAEIN HYPOTHETICAL OXIDOREDUCTASE H11430 pir  H64122 ydfG protein - Haemophilus influenzae (strain Rd KW20) gb AAC23077.1  (U32822) short chain dehydrogenase/reductase [Haemophilus influenzae Rd] Length = 252	Identities = 208/323 (64%), Positives = 249/323 (76%), Gaps = 1/323 (0%) gb AAK05757.1 AE006396_8 (AE006396) phosphate acetyltransferase (EC 2.3.1.8) [Lactococcus lactis subsp. lactis] Length = 326	Identities = 122/204 (59%), Positives = 156/204 (75%) pir  B69878 guanylate kinase homolog yloD - Bacillus subtilis emb CAA74271.1  (Y13937) putative Gmk protein [Bacillus subtilis] emb CAB13441.1  (Z99112) similar to guanylate kinase [Bacillus subtilis] Length = 244
69	79	29	70	72
Contig138 (46516-47259 m)	Contig135 (76527-78527 p)	Contig135 (75631-76392 p)	Contig135 (74577-75569 p)	Contig138 (26798-27427 m)
SA-251.1	SA-2510.2	SA-2511.1	SA-2512.2	SA-2513.2
SeqiD 1508	SeqID 1509	SeqID 1510	SeqID 1511	SeqID 1512

Contig138 (27594-28298 m)  33 Methanobacterium thermoautotrophicum (strain Delta H) gb[AAB85965.1  (AE000909) unknown [Methanobacterium thermoautotrophicum] Length = 188	Identities = 121/199 (60%), Positives = 157/199 (78%), Gaps = 3/199 (1%) emb CAB54585.1  (AJ006400) response regulator [Streptococcus pneumoniae]	Identities = 109/221 (49%), Positives = 150/221 (67%), Gaps = 9/221 (4%) gb AAK02817.1  (AE006110) unknown [Pasteurella multocida]	Identities = 59/131 (45%), Positives = 86/131 (65%), Gaps = 2/131 (1%) sp P44638 LGUL_HAEIN LACTOYLGLUTATHIONE LYASE (METHYLGLYOXALASE) (ALDOKETOMUTASE) (GLYOXALASE I) (GLX I) (KETONE-ALDEHYDE MUTASE) (S-D-LACTOYLGLUTATHIONE METHYLGLYOXAL LYASE) pir   64147   lactoylglutathione   yase (EC 4.4.1.5) - Haemophilus influenzae gb AAC21986.1  (U32717)   lactoylglutathione   yase (gloA) [Haemophilus influenzae Rd] Length = 135	Identities = 139/308 (45%), Positives = 202/308 (65%), Gaps = 3/308 (0%) splQ45539 CSBB_BACSU CSBB PROTEIN pir  JC5173 stress response protein csbB - Bacillus subtilis gb AAB38429.1  (L77099) 44 identity over 302 residues with hypothetical protein from Synechocystis sp, accession hypothetical protein from Synechocystis sp, accession 564 D64006_CD; expression induced by environmental stress; some similarity to glycosyl transferases; two potential membrane-spanning helices [Bacillus subtilis emblCAB12688.1  (299108) stress response protein [Bacillus
-	Contig138 (28348-28938 m) 78	Contig68 (1532-2206 p) 58	Contig68 (2294-2686 p) 62	Contig68 (2815-3741 p) 64
SA-2514.1	SA-2515.2	SA-2516.2	SA-2517.1	SA-2518.1
SeqID 1513	SeqID 1514	SeqID 1515	SeqID 1516	SeqID 1517



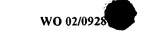
SA-2519.1	Contig68 (3870-4025 p)	28	Identities = 30/46 (65%), Positives = 37/46 (80%) sp P45495 YPEV_LACDL HYPOTHETICAL TRANSPORT PROTEIN IN PEPV 3 REGION (ORF2) pir  S57903 probable amino acid permease - Lactobacillus delbrueckii (fragment) emb CAA83253.1  (231377) potential amino acid permease [Lactobacillus delbrueckii] Length = 175
SA-252.1	Contig138 (47259-48047 m)	209	Identities = 90/224 (40%), Positives = 137/224 (60%), Gaps = 10/224 (4%) splP54953 YXEN_BACSU PROBABLE AMINO-ACID ABC TRANSPORTER PERMEASE PROTEIN YXEN pir  H70075 amino acid ABC transporter (permease) homologyxeN - Bacillus subtilis dbj BAA08330.1  (D45912) homologous to Gln transport system permease proteins [Bacillus subtilis] emb CAB15985.1  (Z99124) similar to amino acid ABC transporter (permease) [Bacillus subtilis]
SA-2520.2	Contig115 (6665-8494 p)	13	Identities = 43/206 (20%), Positives = 91/206 (43%), Gaps = 12/206 (5%) pir] H64496 hypothetical protein MJ1577 - Methanococcus jannaschii gb AAB99606.1  (U67598) M. jannaschii predicted coding region MJ1577 [Methanococcus jannaschii] Length = 598
SA-2522.2	Contig69 (1649-1921 m)	78	Identities = 81/90 (90%), Positives = 85/90 (94%)   gb AAK05667.1 AE006387_10 (AE006387) 30S ribosomal   protein S16 [Lactococcus lactis subsp.   lactis]   Length
SA-2523.2	Contig69 (1397-1639 m)	75	Identities = 39/75 (52%), Positives = 63/75 (84%)   gb AAK05666.1 AE006387_9 (AE006387) UNKNOWN PROTEIN   Lactococcus lactis subsp. lactis   Length = 79
SA-2524.1	Contig69 (3-1265 m)	46	Identities = 115/437 (26%), Positives = 201/437 (45%), Gaps = 26/437 (5%) gb AAC97147.1  (U49397) Nra [Streptococcus pyogenes]

SeqID 1524	SA-2525.1	Contig60 (87-1895 m)	68	Identities = 463/603 (76%), Positives = 540/603 (88%) splP37949 LEPA_BACSU GTP-BINDING PROTEIN LEPA pir  G69649 GTP-binding protein lepA - Bacillus subtilis emb CAA62842.1  (X91655) lepA [Bacillus subtilis] dbj BAA12460.1  (D84432) YqeQ [Bacillus subtilis] emb CAB14493.1  (Z99117) GTP-binding protein [Bacillus cubtilis]
SeqID 1525	SA-2526.1	Contig61 (969-2801 m)	29	), Posit 37.1  (/ s haloc
SeqID 1526	SA-2528.1	Contig61 (82-843 p)	08	Identities = 217/232 (93%), Positives = 226/232 (96%) emb CAB90834.1  (AJ250837) putative transposase [Streptococcus dysgalactiae] Length = 259
SeqID 1527	SA-253.1	Contig138 (48157-48969 m)	49	Identities = 80/273 (29%), Positives = 136/273 (49%), Gaps = 27/273 (9%) emb CAA68052.1  (X99716) collagen binding protein [Lactobacillus reuteri]
SeqID 1528	SA-2531.2	Contig136 (1-1152 p)	26	Identities = 51/228 (22%), Positives = 100/228 (43%), Gaps = 36/228 (15%) emb CAB39029.1  (AL034559) hypothetical protein, PFC0905c [Plasmodium falciparum] Length = 3085
SeqID 1529	SA-2532.1	Contig137 (4715-4921 p)	No Hits found	
SeqID 1530	SA-2533.1	Contig139 (122098-122286 p)	41	Identities = 18/34 (52%), Positives = 26/34 (75%)   gb AAC98435.1  (L29324) unknown [Streptococcus pneumoniae]   Length = 121
SeqID 1531	SA-2534.1	Contig139 (122022-122204 p)	63	Identities = 32/53 (60%), Positives = 39/53 (73%)   gb AAC98435.1  (L29324) unknown [Streptococcus pneumoniae]   Length = 121
SeqID 1532	SA-2535.1	Contig139 (51975-52250 p)	88	Identities = 85/91 (93%), Positives = 88/91 (96%) gb AAD40808.1 L38946_1 (L38946) histone-like DNA-binding protein [Streptococcus pyogenes] Length = 91



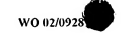
Identities = 126/258 (48%), Positives = 175/258 (66%), Gaps = 2/258 (0%) splO66126 ISPA_MICLU GERANYLTRANSTRANSFERASE (FARNESYL-DIPHOSPHATE SYNTHASE) (FPP SYNTHASE) diphosphate synthase [Micrococcus luteus] Length = 291	Identities = 22/77 (28%), Positives = 41/77 (52%), Gaps = 4/77 (5%) pirj[G69865 hypothetical protein ykuJ - Bacillus subtilis emb[CAA10873.1  (AJ222587) YkuJ protein [Bacillus subtilis] emb[CAB13283.1  (299111) ykuJ [Bacillus subtilis] Length = 79	pund	pund	pund	punc	punc	puno	punc	puno	Identities = 215/439 (48%), Positives = 311/439 (69%), Gaps = 5/439 (1%) splP54475 YQFR_BACSU PROBABLE RNA HELICASE IN CCCA-SODA INTERGENIC REGION pir  D69954 ATP-dependent RNA helicase homolog yqfR - Bacillus subtilis dbj BAA12495.1  (D84432) YqfR (Bacillus subtilis) emb CAB14444.1  (Z99116) similar to ATP-dependent RNA helicase [Bacillus subtilis] Length = 438	pund	punc	pund	Identities = 24/75 (32%), Positives = 46/75 (61%) sp O32233 SECG_BACSU PROBABLE PROTEIN-EXPORT MEMBRANE PROTEIN SECG pir  A70028 hypothetical protein yvaL - Bacillus subtilis emb CAB15368.1  (299121) yvaL [Bacillus subtilis] Length = 76
	39	No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	29	No Hits found	No Hits found	No Hits found	45
Contig139 (45692-46564 p)	Contig139 (38436-38666 p)	1	Contig138 (95981-96244 p)	Contig133 (27162-27320 m)	Contig132 (25696-25839 m)	Contig 127 (19934-20134 p)	Contig135 (30113-30319 p)	Contig135 (29845-30003 p)	Contig135 (27910-28200 m)	Contig138 (49107-50450 m)	Contig123 (36802-36984 m)	Contig116 (30459-30827 m)	Contig116 (60-398 m)	Contig134 (78113-78304 p)
SA-2536.1	SA-2537.1	SA-2539.1	SA-2540.2	SA-2542.1	SA-2543.1	SA-2545.1	SA-2547.1	SA-2548.1	SA-2549.1	SA-255.1	SA-2551.2	SA-2554.1	SA-2555.1	SA-2556.1
SeqID 1533	SeqID 1534	SeqID 1535	SeqID 1536	SeqID 1537	SeqID 1538	SeqID 1539	SeqID 1540	SeqID 1541	SeqID 1542	SeqID 1543	SeqID 1544	SeqID 1545	SeqID 1546	SeqID 1547

SeqID 1548	SA-2557.1	Contig134 (69629-69823 m)	No Hits found	
SeqID 1549	SA-2558.1	Contig134 (69416-69646 p)	No Hits found	
SeqID 1550	SA-2559.1	Contig139 (132392-132580 m)	29	Identities = 36/46 (78%), Positives = 39/46 (84%) gb AAC38687.1  (AF030361) transposase [Streptococcus pneumoniae] gb AAC38702.1  (AF030364) transposase [Streptococcus pneumoniae]
SeqID 1551	SA-256.1	Contig138 (50548-51558 m)	79	Identities = 221/336 (65%), Positives = 271/336 (79%), Gaps = 11/336 (3%) splQ9ZHA5 MRAY_STRPN PHOSPHO-N-ACETYLMURAMOYL-PENTAPEPTIDE-TRANSFERASE (UDP-MURNAC-PENTAPEPTIDE PHOSPHOTRANSFERASE) gblAAC95457.1  (AF068903) undecaprenyl-phosphate-UDP-MurNAc-pentapeptide phospho-MurNAc-pentapeptide transferase [Streptococcus pneumoniae] Length = 326
SeqID 1552	SA-2560.1	Contig139 (133610-133768 p)	37	Identities = 23/35 (65%), Positives = 28/35 (79%) gb AAG38044.1 AF295925_9 (AF295925) Orf28 [Streptococcus pneumoniae] Length = 371
SeqID 1553	SA-2561.1	Contig139 (154195-154389 p)	No Hits found	
SeqID 1554	SA-2562.1	Contig 125 (24635-24835 p)	No Hits found	
SeqID 1555	SA-2564.1	Contig112 (19746-19868 p)	47	Identities = 22/40 (55%), Positives = 27/40 (67%), Gaps = 1/40 (2%) dbj BAB04083.1  (AP001508) transcriptional regulator [Bacillus halodurans] Length = 66
SeqID 1556	SA-2565.1	Contig112 (19523-19741 p)	No Hits found	
SeqID 1557	SA-2566.1		No Hits found	
SeqID 1558	SA-2567.1	Contig111 (13119-13307 p)	No Hits found	
SeqID 1559	SA-257.1	Contig138 (51560-53821 m)	73	Identities = 405/741 (54%), Positives = 556/741 (74%), Gaps = 10/741 (1%) gb AAF17266.1 AF210756_1 (AF210756) penicillinbinding protein 2X [Streptococcus pneumoniae] Length = 750
SeqID 1560	SA-2571.1	Contig106 (21708-21950 p)	41	Identities = 32/83 (38%), Positives = 46/83 (54%) splP55661 Y4TG_RHISN PROBABLE AMINO-ACID ABC TRANSPORTER PERMEASE PROTEIN Y4TG gblAAB91860.1  (AE000098) Y4tG [Rhizobium sp. NGR234] Length = 231



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Identities = 79/110 (71%), Positives = 91/110 (81%), Gaps = 1/110 (0%) splP16680 PHNA_ECOLI PHNA PROTEIN pir  B35718 alkylphosphonate uptake protein phnA [imported] - Escherichia coli gb AAA24337.1  (J05260) phnA protein [Escherichia coli] gb AAA97007.1  (U14003) phnA gene product [Escherichia coli] gb AAC77069.1  (AE000483) orf, hypothetical protein [Escherichia coli] gb AAC77069.1  (AE000483) orf, hypothetical		Identities = 190/228 (83%), Positives = 204/228 (89%)     gb AAK06098.1 AE006430_3 (AE006430) 50S ribosomal protein	Identities = 35/72 (48%), Positives = 42/72 (57%) pir  S59084 hypothetical protein 29.1 - red alga (Chondrus crispus) mitochondrion emb CAA87600.1  (Z47547) unique orf [Chondrus crispus] Length = 79	Identities = 44/99 (44%), Positives = 71/99 (71%) emb CAB01928.1  (Z79691) FtsL [Streptococcus pneumoniae] gb AAC95455.1  (AF068903) YIID [Streptococcus pneumoniae] Length = 105	Identities = 27/42 (64%), Positives = 35/42 (83%)   gb AAD09220.1  (U74080) unknown [Streptococcus gordonii]   Length = 50	Identities = 51/145 (35%), Positives = 91/145 (62%), Gaps = 3/145 (2%) pir  H70091 hypothetical protein yydJ - Bacillus subtilis dbj  BAA11272.1  (D78193) yydJ [Bacillus subtilis] emb  CAB16051.1  (299124) yydJ [Bacillus subtilis]	Identities = 47/106 (44%), Positives = 71/106 (66%) pir  H70091 hypothetical protein yydJ - Bacillus subtilis dbj BAA11272.1  (D78193) yydJ [Bacillus subtilis] emb CAB16051.1  (299124) yydJ [Bacillus subtilis] Length = 240
62	No Hits found No Hits found	88	42	50	53	62	. 09
Contig106 (21244-21573 p)	Contig135 (83485-83640 p) Contig125 (16287-16571 m)	Contig134 (50206-50895 p)	Contig135 (69410-69703 m)	Contig138 (53822-54148 m)	Contig89 (20-172 m)	Contig133 (74342-74776 m)	Contig133 (74719-75069 m)
SA-2572.1	SA-2574.1 SA-2575.1	SA-2576.2	SA-2577.1	SA-258.1	SA-2582.1	SA-2583.1	SA-2584.1
SeqID 1561	SeqID 1562 SeqID 1563	SeqID 1564	SeqID 1565	SeqID 1566	Seq1D 1567	SeqID 1568	SeqID 1569

	100000	Identities = 265/316 (83%), Positives = 288/316 (90%) gb AAC95454.1  (AF068903) YIIC [Streptococcus pneumoniae] Length = 316			Identities = 117/217 (53%), Positives = 168/217 (76%) sp[P42399]YCKA_BACSU PROBABLE AMINO-ACID ABC TRANSPORTER PERMEASE PROTEIN YCKA pir  C69760 amino acid ABC transporter (permease) homology yckA - Bacillus subtilis dbj  BAA06425.1  (D30762) homologue of glutamine permease of H. influenzae [Bacillus subtilis] abj  BAA08971.1  (D50453) homologue of glutamine permease of H. inflenzae [Bacillus subtilis] amb  CAB12131.1  (Z99105) similar to amino acid ABC transporter (permease) [Bacillus subtilis] Length = 226	Identities = 127/276 (46%), Positives = 183/276 (66%), Gaps = 12/276 (4%) splP42400 YCKB_BACSU PROBABLE ABC TRANSPORTER EXTRACELLULAR BINDING PROTEIN YCKB PRECURSOR (ORF2) pirl D69760 amino acid ABC transporter (binding protein) homolog yckB - Bacillus subtilis emb CAB12132.1  (299105) similar to amino acid ABC transporter (binding protein) [Bacillus subtilis] Length = 287
No Hits found	No Hits found	84 gbl	No Hits found	No Hits found	S TE amir 75 St dbjlf H. in s	Ide TRA TRA PRE trans
No H	No Hi		N N	No H		
Contig80 (961-1092 m)	Contig96 (1977-2162 p)	Contig138 (54163-55110 m)	Contig65 (2145-2339 p)	Contig62 (165-344 p)	Contig91 (939-1604 p)	Contig91 (58-915 p)
SA-2586.1	SA-2588.2	SA-259.1	SA-2593.1	SA-2594.1	SA-2596.1	SA-2597.1
SeqID 1570	SeqID 1571	SeqID 1572	SeqID 1573	SeqID 1574	SeqID 1575	SeqID 1576



Identities = 109/218 (50%). Positives = 146/218 (66%), Gaps = 15/218 (6%) splP46338jYOGG_BACSU PROBABLE ABC TRANSPORTER BINDING PROTEIN YOGG PRECURSOR pir[JA69956 phosphate ABC transporter (binding protein) homolog yqgG - Bacıllus subtilis abjiBAA09581.1 (D84432) YqgG Bacıllus subtilis abjiBAA12510.1 (D84432) YqgG [Bacıllus subtilis] emb[CAB14429.1 (299116) alternate gene name yzmB-similar to phosphate ABC transporter (binding protein) [Bacillus subtilis] Length = 300	Identities = 153/445 (34%), Positives = 250/445 (55%); Gaps = 11/445 (2%) emb CAB61253.1  (AJ250422) ORFC [Oenococcus oeni] Length = 463	pun	Identities = 303/417 (72%), Positives = 356/417 (84%), Gaps = 1/417 (0%) splP96489JPROA_STRTR GAMMA-GLUTAMYLPHOSPHATE REDUCTASE (GPR) (GLUTAMATE-5-SEMIALDEHYDE DEHYDROGENASE) (GLUTAMYLGAMMA-SEMIALDEHYDE DEHYDROGENASE) emb[CAA63148.1] (X92418) gamma-glutamyl phosphate reductase [Streptococcus thermophilus] Length = 416		Identities = 63/15 (54%), Positives = 83/115 (71%), Gaps = 1/115 (0%) sp O35046 YOCD_BACSU HYPOTHETICAL 36.4 KDA PROTEIN IN CSAA-DES INTERGENIC REGION pir  A69901 immunity to bacteriotoxins homolog yocD - Bacillus subtilis gb AAB84435.1  (AF027868) YocD [Bacillus subtilis] emb CAB13809.1  (299114) similar to immunity to bacteriotoxins [Bacillus subtilis]	pun
67	54	No Hits found	84	No Hits found	20	No Hits found
Contig104 (8256-8870 p)	Contig135 (79561-80901 p)	Contig137 (15639-15896 p)	Contig138 (55193-56446 m)	Contig110 (3263-3451 m)	Contig110 (3909-4259 p)	Contig111 (11745-11921 p)
SA-2598.1	SA-2599.1	SA-26.1	SA-260.1	SA-2600.1	SA-2602.1	SA-2603.1
SeqiD 1577	SeqID 1578	SeqID 1579	SeqID 1580	SeqID 1581	SeqiD 1582	SeqID 1583

SeqID 1584	SA-2604.1	Contig116 (30551-30826 m)	34	Identities = 30/51 (58%), Positives = 32/51 (61%) pirl F71245 hypothetical protein PHS004 - Pyrococcus horikoshii dbj BAA29293.1  (AP000001) 58aa long hypothetical protein [Pyrococcus horikoshii] Length = 58
SeqID 1585	SA-2605.1	Contig116 (7428-7886 p)	08	Identities = 122/123 (99%), Positives = 123/123 (99%) gb AAG09968.1 AF248037_3 (AF248037) cation efflux system protein [Streptococcus agalactiae]
SeqID 1586	SA-2606.1	Contig 138 (96371-96607 p)	No Hits found	
SeqID 1587	SA-2608.1	Contig139 (135435-135725 m)	No Hits found	
SeqID 1588	SA-2609.1	Contig139 (132781-132939 m)	No Hits found	
SeqID 1589	SA-261.1	Contig138 (56456-57259 m)	98	Identities = 200/265 (75%), Positives = 235/265 (88%) splP96488IPROB_STRTR GLUTAMATE 5-KINASE (GAMMA-GLUTAMYL KINASE) (GK) emb CAA63147.1  (X92418) gamma-glutamyl kinase [Streptococcus thermophilus] Length = 267
SeqID 1590	SA-2610.1	Contig139 (132368-132517 p)	09	Identities = 19/36 (52%), Positives = 30/36 (82%) pirj T50042 hypothetical protein tasA [imported] - Streptococcus pneumoniae emb CAA59773.1  (X85787) tasA [Streptococcus pneumoniae] Length = 359
SeqID 1591	SA-2612.1	Contig139 (93569-93832 p)	No Hits found	
SeqID 1592	SA-2615.1	Contig139 (52350-52646 m)	No Hits found	
SeqID 1593	SA-2618.1	Contig139 (18458-18772 m)	35	Identities = 30/71 (42%), Positives = 37/71 (51%), Gaps = 1/71 (1%) pirj[G72514 hypothetical protein APE2092 - Aeropyrum pernix (strain K1) dbj[BAA81103.1] (AP000063) 101aa long hypothetical protein [Aeropyrum pernix] Length = 101
SeqID 1594	SA-2620.1	Contig138 (81930-82160 m)	No Hits found	

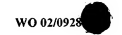
SeqID 1595 SeqID 1596 SeqID 1596 SeqID 1599 SeqID 1600 SeqID 1601 SeqID 1603 SeqID 1603 SeqID 1603 SeqID 1605	SA-263.1 SA-2632.1 SA-2635.1 SA-264.1 SA-264.1 SA-264.1 SA-265.1 SA-265.1 SA-265.1	Contig138 (57400-58785 p)  Contig137 (2699-2956 p)  Contig136 (45825-45980 p)  Contig138 (5893-60597 m)  Contig138 (29120-29389 m)  Contig134 (73407-73562 m)  Contig133 (60601-61335 m)  Contig133 (68964-69245 p)  Contig133 (60101-60367 p)  Contig132 (48100-48423 m)	No Hits found	Identities = 174/47 (38%), Positives = 267/447 (58%), Gaps = 10/447 (2%) pir  T47097 hypothetical protein N17C (imported] - Bacillus subtilis dbj BAA06562.1  (D29865) hypothetical protein [Bacillus subtilis] dbj BAA06256.1  (D29865) hypothetical 64.7-kDa protein [Bacillus subtilis] mb[CAB15963.1  (299124) phosphotransferase system (PTS) beta-glucoside-specific enzyme IIABC component [Bacillus subtilis] Length = 609
SeqID 1606	SA-2655.1	Contig132 (48095-48463 p)	89	Identities = 116/122 (95%), Positives = 120/122 (98%) splQ9WVZ2 RL14_STRPN 50S RIBOSOMAL PROTEIN L14 gb AAD33266.1 AF126059_7 (AF126059) RpL14 (Streptococcus pneumoniae] gb AAD33275.1  (AF126060) RpL14 (Streptococcus pneumoniae] gb AAD33284.1  (AF126061) RpL14 (Streptococcus pneumoniae] gb AAD33284.1  (AF126061)

puno				puno	puno	Identities = 403/661 (60%). Positives = 520/661 (77%), Gaps =   8/661 (1%) dbj BAB06071.1  (AP001515) transketolase [Bacillus halodurans]	puno,	puno	Identities = 40/83 (48%), Positives = 64/83 (76%) pir  A69742 conserved hypothetical protein yazA - Bacillus subtilis emb CAB11811.1  (299104) similar to hypothetical proteins [Bacillus subtilis] Length = 99	Identities = 48/191 (25%), Positives = 88/191 (45%), Gaps = 9/191 (4%) emb CAB65412.1  (AJ243530) putative Mga-like regulatory protein [Streptococcus dysgalactiae subsp. dysgalactiae]	puno	puno	puno	Identities = 25/72 (34%), Positives = 36/72 (49%) pir  G72510 hypothetical protein APE2061 - Aeropyrum pernix (strain K1) dbj BAA81071.1  (AP000063) 114aa long hypothetical protein [Aeropyrum pernix] Length = 114	Identities = 105/423 (24%), Positives = 198/423 (45%), Gaps = 15/423 (3%) splP37061 NAOX_ENTFA NADH OXIDASE (NOXASE) pir  S26965 NADH oxidase - Enterococcus faecalis emb CAA48728.1  (X68847) NADH oxidase [Enterococcus faecalis]	
No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	No Hits found	77	No Hits found	No Hits found	53	18	No Hits found	No Hits found	No Hits found	33	44	
Contig132 (25921-26223 p)		Contig131 (2878-3129 m)	Contig 130 (70425-70709 p)	Contig130 (38146-38331 m)	Contig130 (27964-28215 p)	Contig138 (61873-63858 m)	Contig129 (57906-58148 m)	Contig129 (47951-48133 p)	Contig129 (21896-22162 p)	Contig138 (63983-65413 m)	Contig126 (41736-42008 p)	Contig126 (33868-34071 p)	Contig125 (36869-37171 m)	Contig124 (23609-23926 p)	Contig138 (65404-66741 m)	
SA-2657.1	SA-266.1	SA-2664.1	SA-2666.1	SA-2668.1	SA-2669.1	SA-267.1	SA-2670.1	SA-2672.1	SA-2673.1	SA-268.1	SA-2681.1	SA-2683.1	SA-2687.1	SA-2689.1	SA-269.1	
SeqID 1607	SeqID 1608	SeqID 1609	SeqID 1610	SeqID 1611	SeqID 1612	SeqID 1613	SeqID 1614	SeqID 1615	SeqID 1616	SeqID 1617	SeqID 1618	SeqID 1619	SeqID 1620	SeqID 1621	SeqID 1622	-

SeqID 1624	SA-2691.1	Contig123 (39313-39621 p)	36	Identities = 21/61 (34%), Positives = 38/61 (61%), Gaps = 1/61 (1%) ref[NP_076769.1] Orf18 [bacteriophage blL310] gb AAK04132.1 AE006242_11 (AE006242) prophage ps1 protein 10 [Lactococcus lactis subsp. lactis] gb AAK08422.1 AF323671_18 (AF323671) Orf18 [bacteriophage blL310] Length = 93
SeqID 1625	SA-2696.1	Contig122 (25328-25630 m)	34	Identities = 22/56 (39%), Positives = 35/56 (62%), Gaps = 4/56 (7%) splP36417 GBF_DICDI G-BOX BINDING FACTOR (GBF) pir  A53185 G-box-binding factor - slime mold (Dictyostelium discoideum) gb AAA21021.1  (L29075) G-box binding factor   [Dictyostelium discoideum] Length = 708
SeqID 1626	SA-270.1	Contig138 (66829-67542 m)	18	Identities = 150/230 (65%), Positives = 194/230 (84%), Gaps = 1/230 (0%) splP52281 GLPF_STRPN GLYCEROL UPTAKE FACILITATOR PROTEIN pir  S67937 glycerol uptake facilitator GlpF - Streptococcus pneumoniae (strain P13) gb AAA91618.1  (U12567) glycerol uptake facilitator [Streptococcus pneumoniae] Length = 233
SeqID 1627	SA-2703.1	Contig119 (308-1129 m)	50	Identities = 94/304 (30%), Positives = 152/304 (49%), Gaps = 32/304 (10%) gbjAAK04228.1jAE006251_4 (AE006251) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 304
SeqID 1628	SA-2705.1	Contig118 (27339-27527 p)	No Hits found	
SeqID 1629	SA-2707.1	Contig118 (17969-18169 p)	80	Identities = 52/66 (78%), Positives = 60/66 (90%)   gb AAK05941.1 AE006414_7 (AE006414) 50S ribosomal protein   L35 [Lactococcus lactis subsp.   lactis]
SeqID 1630	SA-2709.1	Contig118 (17399-17929 p)	74	Identities = 112/169 (66%), Positives = 134/169 (79%) sp O53084 IF3_LISMO TRANSLATION INITIATION FACTOR IF- 3 emb CAA68920.1  (Y07640) translation initiation factor, IF3 [Listeria monocytogenes] Length = 171
SeqID 1631	SA-271.1	Contig138 (67539-69368 m)	87	Identities = 464/608 (76%), Positives = 539/608 (88%) gb AAC34740.1  (U94770) alpha-glycerophosphate oxidase [Streptococcus pneumoniae] Length = 608

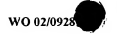


SeqID 1632	SA-2710.1	Contig118 (16555-17238 p)	73	Identities = 139/223 (62%), Positives = 171/223 (76%), Gaps = 3/223 (1%) gb]AAK05801.1 AE006401_2 (AE006401) cytidine monophosphate kinase (EC 2.7.4.14) [Lactococcus lactis subsp. lactis]
SeqID 1633	SA-2711.1	Contig118 (16244-16408 m)	No Hits found	
SeqID 1634	SA-2712.1	Contig118 (16020-16544 p)	42	Identities = 46/167 (27%), Positives = 82/167 (48%), Gaps = 15/167 (8%) gb AAK05013.1 AE006326_4 (AE006326) teichoic acid ABC transporter ATP binding protein [Lactococcus lactis subsp. lactis] Length = 466
SeqID 1635	SA-2715.1	Contig117 (11061-11336 p)	No Hits found	
SeqID 1636	SA-2718.1	Contig116 (21405-21707 p)	No Hits found	
SeqID 1637	SA-2719.1	Contig116 (19624-19911 p)	No Hits found	
SeqID 1638	SA-272.1	Contig138 (69381-70889 m)	82	Identities = 383/501 (76%), Positives = 442/501 (87%), Gaps = 1/501 (0%) spj034154jGLPK_ENTFA GLYCEROL KINASE (ATP:GLYCEROL 3-PHOSPHOTRANSFERASE) (GLYCEROKINASE) (GK) gbjAAB69986.1j (U94356) glycerol kinase [Enterococcus faecalis] Length = 501
SeqID 1639	SA-2722.1	Contig115 (26094-26351 m)	45	Identities = 36/58 (62%), Positives = 39/58 (67%) gb AAG12204.1 AF287482_5 (AF287482) Orf122 [Chlorobium tepidum] Length = 121
SeqID 1640	SA-2726.1	Contig113 (25586-25693 p)	No Hits found	
SeqiD 1641	SA-273.1	Contig138 (71003-71266 m)	43	Identities = 36/79 (45%), Positives = 48/79 (60%), Gaps = 1/79 (1%) pir  G83401 hypothetical protein PA1960 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG05348.1 AE004622_10 (AE004622) hypothetical protein [Pseudomonas aeruginosa] Length = 240
SeqID 1642	SA-2733.1	Contig108 (18783-18911 p)	No Hits found	
SeqID 1643	SA-274.1	Contig138 (71355-71612 m)	28	Identities = 41/72 (56%), Positives = 56/72 (76%) pir  E69894 hypothetical protein ynzC - Bacillus subtilis emb  CAB13672.1  (Z99113) ynzC (Bacillus subtilis) Length = 77
SeqID 1644	SA-2741.1	Contig 102 (13042-13308 p)	No Hits found	
SeqID 1645	SA-2747.1	Contig98 (8006-8278 p)	No Hits found	



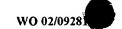
A-275.1 -2752.1 -2756.1 -2758.1 -2758.1 -2760.1 -2761.1	Identities = 315/687 (45%), Positives = 445/687 (63%), Gaps = 21/687 (3%) splP54381 SYGB_BACSU GLYCYL-TRNA SYNTHETASE BETA CHAIN (GLYCINETRNA LIGASE BETA CHAIN (GLYCINETRNA LIGASE BETA CHAIN (GLYRS) pir[ B69636 glycinetRNA ligase (EC 6.1.1.14) beta chain glyS - Bacillus subtilis ability (71624-73663 m)	Contig94 (10312-10587 p) No Hits found	dentities = 32/79 (40%), Positives = 48/79 (60%)   gb AAK05381.1 AE006360_10 (AE006360) HYPOTHETICAL     PROTEIN [Lactococcus lactis subsp.   lactis] Length = 79	Contig91 (6359-6622 m) No Hits found	Contig91 (5806-6126 m) No Hits found	Contig91 (5708-6319 p)  46  2/153 (1%) gb AAK05259.1 AE006348_4 (AE006348) UNKNOWN PROTEIN [Lactococcus lactis subsp. lactis] Length = 199	Identities = 64/176 (36%), Positives = 99/176 (55%), Gaps = 67/176 (36%), Positives = 99/176 (55%), Gaps = 67/16 (36%), Positives = 99/176 (55%), Gaps = 67/176 (36%), Positives = 99/176 (36%), Positives = 99/176 (36%), Positives = 99/176 (36%), Gaps = 67/176 (36%), Positives = 99/176 (36%), Positive	Identities = 33/106 (31%), Positives = 53/106 (49%), Gaps = 8/106 (13-960 m)	Contig90 (10903-11157 m) No Hits found	Identities = 298/298 (100%), Positives = 298/298 (100%)   gb AAK14387.1 AF338416_1 (AF338416) glyceraldehyde 3-phosphate dehydrogenase [Streptococcus agalactiae]	
	SA-275.1	SA-2752.1	SA-2756.1	SA-2757.1	SA-2758.1	SA-2759.1	SA-276.1	SA-2760.1	SA-2761.1	SA-2763.1	

Identities = 79/142 (55%), Positives = 102/142 (71%), Gaps = 5/142 (3%) gb AAK05046.1 AE006329_5 (AE006329) UNKNOWN PROTEIN [Lactococcus lactis subsp. lactis] Length = 159	Identities = 38/131 (29%), Positives = 70/131 (53%), Gaps = 5/131 (3%) pir  B83475 hypothetical protein PA1353 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG04742.1 AE004565_1 (AE004565) hypothetical protein [Pseudomonas aeruginosa] Length = 137		Identities = 221/287 (77%), Positives = 250/287 (87%) splQ9KD49 SYGA_BACHD GLYCYL-TRNA SYNTHETASE ALPHA CHAIN (GLYCINETRNA LIGASE ALPHA CHAIN) (GLYRS) dbj BAB05089.1  (AP001511) glycyl-tRNA synthetase (alpha subunit) [Bacillus halodurans] Length = 297	Identities = 17/58 (29%), Positives = 27/58 (46%), Gaps = 1/58 (1%) ref[NP_042981.1  U88 [Human herpesvirus 6] emb[CAA58337.1  (X83413) U88 [Human herpesvirus 6] Length = 413			Identitles = 26/51 (50%), Positives = 37/51 (71%), Gaps = 2/51 (3%) gb AAK04732.1 AE006296_6 (AE006296) DNA-directed RNA polymerase delta chain (EC 2.7.7.6) [Lactococcus lactis subsp. lactis] Length = 187		Identities = 71/168 (42%), Positives = 105/168 (62%) pir  F81147 probable integral membrane protein NMA1102 [Imported] - Neisseria meningitidis (group B strain MD58, group A strain Z2491) gb AAF41294.1  (AE002440) conserved hypothetical protein [Neisseria meningitidis MC58] emb CAB84365.1  (AL162755) putative integral membrane protein [Neisseria meningitidis Z2491] Length = 169
29	50	No Hits found	72	24	No Hits found	No Hits found	64	No Hits found	57
Contig84 (10458-10913 m)	Contig84 (10038-10448 m)	Contig83 (5598-5870 p)	Contig138 (74308-75225 m)	Contig83 (3561-3887 p)	Contig80 (1912-2166 m)	Contig80 (902-1168 p)	Contig78 (6209-6433 p)	Contig69 (2000-2224 p)	Contig138 (75535-76077 m)
SA-2767.1	SA-2768.1	SA-2769.1	SA-277.1	SA-2770.1	SA-2771.1	SA-2772.1	SA-2773.1	SA-2776.1	SA-278.2
SeqID 1657	SeqID 1658	SeqID 1659	SeqID 1660	SeqID 1661	SeqID 1662	SeqID 1663	SeqID 1664	SeqID 1665	SeqID 1666



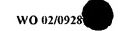
Seq1D 1667	SA-279.1	Contig126 (42614-42796 p)	No Hits found	
SeqID 1668	SA-28.1	Contig137 (13147-15513 p)	20	Identities = 96/354 (27%), Positives = 161/354 (45%), Gaps = 61/354 (17%) ref[NP_053238.1  pXO2-84 [Bacillus anthracis] gb AAF13688.1 AF188935_86 (AF188935) pXO2-84 [Bacillus anthracis] Length = 490
SeqID 1669	SA-280.1	Contig126 (42553-46098 m)	69	Identities = 629/1241 (50%), Positives = 825/1241 (65%), Gaps = 75/1241 (6%) gb AAG33958.1 AF217414_1 (AF217414)   pullulanase [Streptococcus pneumoniae]   Length = 1287
SeqiD 1670	SA-282.1	Contig126 (41501-42421 m)	61	Identities = 134/299 (44%), Positives = 197/299 (65%) dbj BAB08178.1  (AB036768) exfoliative toxin A [Staphylococcus hyicus] Length = 306
SeqID 1671	SA-283.1	Contig126 (41283-41459 p)	No Hits found	
SeqID 1672	SA-285.1	Contig126 (40282-41184 m)	. 62	Identities = 139/311 (44%), Positives = 200/311 (63%), Gaps = 21/311 (6%) splQ9KAC3 MIAA_BACHD TRNA DELTA(2)-ISOPENTENYLPYROPHOSPHATE TRANSFERASE (IPP TRANSFERASE) (ISOPENTENYL-DIPHOSPHATE: TRNA ISOPENTENYLTRANSFERASE) (IPTASE) (IPPT) dbj BAB06085.1  (AP001515) tRNA isopentenylpyrophosphate transferase [Bacillus halodurans] Length = 314
SeqID 1673	SA-286.1	Contig126 (38953-40191 m)	61	Identities = 183/406 (45%), Positives = 255/406 (62%), Gaps = 12/406 (2%) dbj BAB06081.1  (AP001515) unknown conserved protein [Bacillus halodurans] Length = 418
SeqID 1674	SA-287.1	Contig126 (38313-38960 m)	No Hits found	
SeqID 1675	SA-288.1	Contig126 (37342-38271 m)	49	Identities = 156/309 (50%), Positives = 209/309 (67%), Gaps = 5/309 (1%) splP54548 YQJK_BACSU HYPOTHETICAL 34.0 KDA PROTEIN IN GLNQ-ANSR INTERGENIC REGION pir  C69964 conserved hypothetical protein yqlK - Bacillus subtilis dbj BAA12617.1  (D84432) YqjK (Bacillus subtilis] emb CAB14316.1  (299116) similar to hypothetical proteins [Bacillus subtilis]
SeqID 1676	SA-29.1	Contig137 (12635-13093 p)	No Hits found	

SeqID 1677	SA-290.1	Contig126 (36579-37340 m)	59	Identities = 100/253 (39%), Positives = 152/253 (59%), Gaps = 2/253 (0%) splP54554 YQJQ_BACSU HYPOTHETICAL OXIDOREDUCTASE IN ANSR-BMRU INTERGENIC REGION pir  A69965 ketoacyl reductase homolog yqjQ - Bacillus subtilis dbj BAA12623.1  (D84432) YqjQ [Bacillus subtilis] emb CAB14310.1  (Z99116) similar to ketoacyl reductase [Bacillus subtilis]
SeqID 1678	SA-291.1	Contig126 (34384-36582 m)	09	Identities = 276/772 (35%), Positives = 447/772 (57%), Gaps = 45/772 (5%) pir  H69980 single-strand DNA-specific exonuclease homolog yrvE - Bacillus subtilis emb CAB14721.1  (299118) similar to single-strand DNA-specific exonuclease [Bacillus subtilis] Length = 786
SeqID 1679	SA-292.1	Contig126 (31668-34187 m)	12	Identities = 65/220 (29%), Positives = 104/220 (46%), Gaps = 33/220 (15%) sp P11701 SACB_STRMU LEVANSUCRASE PRECURSOR (BETA-D-FRUCTOFURANOSYL TRANSFERASE) (SUCROSE 6-FRUCTOSYL TRANSFERASE) pir  B28551 levansucrase (EC 2.4.1.10) precursor - Streptococcus mutans (strain GS-5) gb AAA88584.1  (M18954) fructosyltransferase [Streptococcus mutans] Length = 797
SeqID 1680	SA-293.1	Contig126 (31027-31545 m)	74	Identities = 110/170 (64%), Positives = 135/170 (78%) splO34443 APT_BACSU ADENINE PHOSPHORIBOSYLTRANSFERASE (APRT) plr  B69587 adenine phosphoribosyltransferase (EC 2.4.2.7) - Bacillus subtilis gb AAC46040.1  (U86377) adenine phosphoribosyltransferase; Apt [Bacillus subtilis] emb CAB14720.1  (Z99118) adenine phosphoribosyltransferase [Bacillus subtilis] Length = 170
SeqID 1681	SA-294.1	Contig126 (30229-30909 m)	78	Identities = 140/227 (61%), Positives = 179/227 (78%) dbj BAA11244.1  (D78182) ORF2 [Streptococcus mutans] Length = 231
SeqID 1682	SA-295.1	Contig126 (29442-30125 m)	1.1	Identities = 132/226 (58%), Positives = 168/226 (73%) dbj BAA11245.1  (D78182) ORF3 [Streptococcus mutans] Length = 232



SeqID 1683	SA-296.1	Contig126 (28664-29455 m)	83	Identities = 186/262 (70%), Positives = 224/262 (84%)   dbj BAA11246 1  (D78182) ORF4 (Streptococcus mutans)   Length = 262
SeqID 1684	SA-297.1	Contig126 (27552-28655 m)	55	Identities = 147/368 (39%), Positives = 210/368 (56%), Gaps = 13/368 (3%) spl032159 YURR_BACSU HYPOTHETICAL 39.4 KD OXIDOREDUCTASE IN HOM-MRGA INTERGENIC REGION pir  A70019 opine catabolism homolog yurR - Bacillus subtilis emb CAB15253.1  (299120) similar to opine catabolism [Bacillus subtilis] Length = 372
SeqID 1685	SA-298.3	Contig126 (26639-27493 m)	68	Identities = 247/275 (89%), Positives = 260/275 (93%) gb[AAD19913.1  (AF105113) glucose-1-phosphate thymidylyl transferase [Streptococcus pneumoniae] Length = 289
SeqID 1686	SA-299.3	Contig126 (26032-26625 m)	98	Identities = 162/198 (81%), Positives = 183/198 (91%) gb AAC78675.1  (AF094575) dTDP-4-keto-6-deoxyglucose-3,5- epimerase Cps19aM [Streptococcus pneumoniae] Length = 198
SeqID 1687	SA-3.1	Contig137 (40498-42447 p)	90	Identities = 188/659 (28%), Positives = 327/659 (49%), Gaps = 29/659 (4%) ref NP_053164.1  pXO2.09 [Bacillus anthracis] gb AAF13614.1 AF188935_12 (AF188935) pXO2.09 [Bacillus anthracis] Length = 643
SeqID 1688	SA-30.1	Contig137 (11818-12618 p)	22	Identities = 48/115 (41%), Positives = 64/115 (54%), Gaps = 3/115 (2%) emb CAA59264.1  (X84793) streptodornase [Streptococcus pyogenes] Length = 385
SeqID 1689	SA-301.1	Contig126 (24779-25825 m)	16	Identities = 325/347 (93%), Positives = 340/347 (97%) splP95780JRMLB_STRMU DTDP-GLUCOSE 4,6-DEHYDRATASE dbj BAA11249.1  (D78182) dTDP-glucose-4,6-dehydratase [Streptococcus mutans] Length = 348
SeqID 1690	SA-303.1	Contig 126 (22369-22569 m)	No Hits found	
SeqID 1691	SA-305.1	Contig126 (21509-24727 p)	66	Identities = 1070/1072 (99%), Positives = 1071/1072 (99%)   emb CAA75865.1  (Y15903) hyaluronate lyase [Streptococcus   agalactiae]

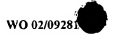
SeqID 1692	SA-306.1	Contig126 (20810-21292 m)	88	Identities = 132/160 (82%), Positives = 146/160 (90%), Gaps = 1/160 (0%) sp P95781 MUTX_STRMU MUTATOR MUTT PROTEIN (7,8-DIHYDRO-8-OXOGUANINE-TRIPHOSPHATASE) (8-OXO-DGTPASE) (PGTP PYROPHOSPHOHYDROLASE) dbj BAA11250.1  (D78182) MutX [Streptococcus mutans] Length = 159
	SA-307.1	Contig126 (20359-20697 m)	45	Identities = 46/150 (30%), Positives = 73/150 (48%), Gaps = 16/150 (10%) dbj BAB07445.1  (AP001519) unknown conserved protein in others [Bacillus halodurans] Length = 152
L	SA-308.1	Contig126 (19077-20267 m)	76	Identities = 243/393 (61%), Positives = 303/393 (76%), Gaps = 8/393 (2%) gb AAD00285.1  (U78604) putative membrane protein [Streptococcus mutans]
-	SA-309.1	Contig126 (17861-19102 m)	49	Identities = 133/347 (38%), Positives = 207/347 (59%), Gaps = 5/347 (1%) gb AAA25160.1  (L16975) ORF1 [Lactococcus lactis] Length = 349
	SA-31.1	Contig137 (11471-11818 p)	No Hits found	
	SA-311.1	Contig126 (16069-17751 m)	82	Identities = 395/559 (70%), Positives = 465/559 (82%), Gaps = 8/559 (1%) gb AAA25161.1  (L16975) alpha-acetolactate synthase [Lactococcus lactis]
	SA-312.1	Contig126 (15336-16055 m)	72	Identities = 139/239 (58%), Positives = 186/239 (77%), Gaps = 3/239 (1%) gb AAB37482.1  (S82499) alpha-acetolactate decarboxylase, AldB [Lactococcus lactis, ssp. lactis, NCDO2118, Peptide, 236 aa] gb AAB81923.1  (U92974) AldB [Lactococcus lactis] Length = 236
	SA-313.2	Contig126 (13627-15282 p)	82	Identities = 393/550 (71%), Positives = 462/550 (83%) emb[CAA46282.1] (X65164) fibronectin-binding protein-like protein A [Streptococcus gordonii] Length = 550



5	Sontig126 (8670-10331 m) 64	
63 No Hits found No Hits found 21	Contig126 (7825-8613 m)  20ntig137 (11161-11478 p)  Contig126 (7025-7684 m)  Contig126 (6333-6878 p)	SA-319.1 Contig126 (7825-8613 m)  SA-32.1 Contig137 (11161-11478 p)  SA-320.1 Contig126 (7025-7684 m)  SA-321.1 Contig126 (6333-6878 p)



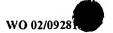
SeqID 1708	SA-322.1	Contig126 (6116-6787 m)	88	Identities = 103/220 (46%), Positives = 141/220 (63%), Gaps = 3/220 (1%) splP72012 RPIA_METTH PROBABLE RIBOSE 5-PHOSPHATE ISOMERASE (PHOSPHORIBOISOMERASE) pir  G69180 ribose 5-phosphate isomerase - Methanobacterium thermoautotrophicum (strain Delta H) dbj BAA13646.1  (D88555) orf2 [Methanobacterium thermoautotrophicum] gb AAB85114.1  (AE000842) ribose 5-phosphate isomerase [Methanobacterium thermoautotrophicum] Length = 226
SeqID 1709	SA-323.1	Contig126 (4848-6059 m)	. 62	Identities = 275/408 (67%), Positives = 325/408 (79%), Gaps = 7/408 (1%) spl032808jDEOB_LACLC PHOSPHOPENTOMUTASE (PHOSPHODEOXYRIBOMUTASE) gb AAC45496.1  (U80410) phosphopentomutase [Lactococcus lactis subsp. cremoris] Length = 411
SeqID 1710	SA-324.1	Contig126 (4390-4800 m)	55	Identities = 73/130 (56%), Positives = 93/130 (71%) splP52147 ARC2_ECOLI ARSENATE REDUCTASE (ARSENICAL PUMP MODIFIER) gblAAB09628.1  (U38947) ArsC [Plasmid R46] Length = 141
SeqID 1711	SA-325.1	Contig126 (3542-4351 m)	74	Identities = 149/272 (54%), Positives = 201/272 (73%), Gaps = 3/272 (1%) dbj BAB05250.1  (AP001512) purine nucleoside phosphorylase [Bacillus halodurans] Length = 272
SeqID 1712	SA-326.1	Contig126 (2284-3540 m)	90	Identities = 121/410 (29%), Positives = 215/410 (51%), Gaps = 19/410 (4%) gblAAD53928.1 AF179611_12 (AF179611) chloride channel protein [Zymomonas mobilis] Length = 425
SeqID 1713	SA-327.1	Contig126 (1590-2300 m)	82	Identities = 177/216 (81%), Positives = 197/216 (90%) splQ56037 DEOD_STRTR PURINE NUCLEOSIDE PHOSPHORYLASE (INOSINE PHOSPHORYLASE) (PNP) gb AAC44007.1  (U40830) DeoD [Streptococcus thermophilus] prf  2209356A deoD gene [Streptococcus thermophilus] Length = 216
SeqID 1714	SA-328.1	Contig126 (814-1581 m)	No Hits found	



	SA-329.1	Contig126 (169-789 p)	66	Identities = 262/262 (100%), Positives = 262/262 (100%) pir  T44638 capsular polysaccharide biosynthesis protein cpsY [imported] - Streptococcus agalactiae emb CAB36980.1  (Y17218) CpsY protein [Streptococcus agalactiae] emb CAB36982.2  (Y17241) CpsY protein [Streptococcus agalactiae] agalactiae] gb AAD53064.1 AF163833_1 (AF163833) CpsY [Streptococcus agalactiae] Length = 307
	SA-330.2	Contig132 (27403-27660 p)	No Hits found	
	SA-331.2	Contig132 (26320-27393 p)	69	Identities = 212/347 (61%), Positives = 261/347 (75%), Gaps = 3/347 (0%) emb[CAA04376.1  (AJ000883) purK [Lactococcus lactis] Length = 349
!	SA-333.1	Contig132 (25842-26333 p)	28	Identities = 102/158 (64%), Positives = 129/158 (81%) emb[CAA04375.1] (AJ000883) purE [Lactococcus lactis] Length = 161
	SA-334.1	Contig132 (24299-25561 p)	99	Identities = 239/419 (57%), Positives = 300/419 (71%), Gaps = 7/419 (1%) splQ9ZF44 PUR2_LACLA PHOSPHORIBOSYLAMINEGLYCINE LIGASE (GARS) (GLYCINAMIDE RIBONUCLEOTIDE SYNTHETASE) (PHOSPHORIBOSYLGLYCINAMIDE SYNTHETASE) emb CAA04374.1  (AJ000883) purD [Lactococcus lactis] Length = 412
	SA-335.1	Contig132 (23212-24015 m)	. 4	Identities = 66/258 (25%), Positives = 119/258 (45%), Gaps = 9/258 (3%) splP26833 YNGB_CLOPE HYPOTHETICAL 31.2 KDA PROTEIN IN NAGH 5 REGION (ORFB) pir  S43902 hypothetical protein B - Clostridium perfringens gb AAA23257.1  (M81878) unknown [Clostridium perfringens] Length = 279
	SA-336.1	Contig132 (22212-23189 p)	55	Identities = 133/299 (44%), Positives = 188/299 (62%), Gaps = 1/299 (0%) pir  S41858 hypothetical protein - Thermoanaerobacterium saccharolyticum Length = 320



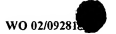
	SA-337.1	Contig132 (21323-22204 p)	37	Identities = 78/160 (48%), Positives = 110/160 (68%)  sp P26832 YNGA_CLOPE HYPOTHETICAL PROTEIN IN NAGH 5 REGION (ORFA) pir  S43901 hypothetical protein A - Clostridium perfringens (fragment) gb AAA23256.1  (M81878) unknown [Clostridium perfringens] Length = 182
S	SA-338.1	Contig132 (20389-21306 p)	58	Identities = 162/225 (72%), Positives = 191/225 (84%) emb CAA69950 1  (Y08695) putative acylneuraminate lyase [Clostridium tertium] Length = 226
"	SA-339.1	Contig132 (19703-20392 p)	32	Identities = 40/148 (27%), Positives = 74/148 (49%), Gaps = 4/148 (2%) dbj BAB05827.1 (AP001514) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 214
	SA-34.1	Contig137 (10492-10767 p)	40	Identities = 30/91 (32%), Positives = 50/91 (53%), Gaps = 6/91 (6%) gb AAF73773.1 AF154006_1 (AF154006) surface protein PspC [Streptococcus pneumoniae] Length = 678
	SA-340.1	Contig132 (19267-19710 p)	43	dentities = 48/151 (31%), Positives = 66/151 (42%), Gaps = 5/151 (3%) gb AAC44392.1  (U43526) ORF-1 [Streptococcus pneumoniae] Length = 150
1	SA-342.1	Contig132 (18424-19254 p)	99	Identities = 94/262 (35%), Positives = 158/262 (59%), Gaps = 1/262 (0%) pir F72379 sugar ABC transporter, permease protein Thermotoga maritima (strain MSB8) gb AAD35515.1 AE001721_14 (AE001721) sugar ABC transporter, permease protein [Thermotoga maritima] Length = 271
·	SA-343.1	Contig132 (17527-18414 p)	51	Identities = 106/289 (36%), Positives = 168/289 (57%), Gaps = 6/289 (2%) dbj BAB05584. I  (AP001513) sugar transport system (permease) (binding protein dependent transporter) [Bacillus halodurans] Length = 309
	SA-344.1	Contig132 (16123-17439 p)	20	Identities = 54/187 (28%), Positives = 90/187 (47%), Gaps = 14/187 (7%) pir[ F69796 sugar-binding protein homolog yesO - Bacillus subtilis emb CAB12516.1  (299107) similar to sugar-binding protein [Bacillus subtilis] Length = 412



Identities = 107/226 (47%), Positives = 148/226 (65%), Gaps = 6/226 (2%) pir[ C70180 conserved hypothetical protein BB0644 - Lyme disease spirochete gb[AAC66999.1] (AE001166) conserved hypothetical protein [Borrelia burgdorferi] Length = 232	Identities = 434/434 (100%), Positives = 434/434 (100%) gb AAG18476.1  (AF151359) group B streptococcal surface immunogenic protein [Streptococcus agalactiae] gb AAG18477.1  (AF151361) group B streptococcal surface immunogenic protein [Streptococcus agalactiae] Length = 434	Identities = 163/274 (59%), Positives = 196/274 (71%), Gaps = 11/274 (4%) gb AAC46072.1  (U50357) zoocin A endopeptidase [Streptococcus zooepidemicus]	Identities = 310/515 (60%), Positives = 390/515 (75%), Gaps = 4/515 (0%) dbj BAB04352.1  (AP001509) phosphoribosylaminoimidazolecarboxamide formyltransferase/IMP cyclohydrolase [Bacillus halodurans] Length = 511	Identities = 65/263 (24%), Positives = 117/263 (43%), Gaps = 34/263 (12%) pir  H69779 antibiotic resistance protein homolog ydfB - Bacillus subtilis dbj BAA19369.1 (AB001488) SIMILAR TO BACILLUS CEREUS ZWITTERMICIN A-RESISTANCE GENE. [Bacillus subtilis] emb CAB12342.1 (299106) similar to intibiotic resistance protein [Bacillus subtilis] Length = 261	Identities = 137/152 (90%), Positives = 140/152 (91%), Gaps = 2/152 (1%) gb AAF08602.1 U70775_1 (U70775)
Identities = 107/2/ 6/226 (2%) pir  C7/ Lyme disease conserved hypothe	Identities = 434/434 (1 gbJAAG18476.1] (AF15: immunogenic protein gbJAAG18477.1] (AF15: immunogenic protein	Identities = 163/27 11/274 (4%) gb AA [Streptococc	Identities = 310/5' 4/515 ( phospho formyltran	Identities = 65/263 (24%), Positive: 34/263 (12%) pir  H69779 antibiotic ydfB - Bacillus subtilis dbj BAA193 TO BACILLUS CEREUS ZWITTE GENE. [Bacillus subtilis] emb CABantibiotic resistance protein [Bacillus = 261	Identities = 137/18 2/152 (1%)
62	26	62	74	44	61
Contig132 (15378-16076 p)	Contig132 (13827-15131 p)	Contig132 (12781-13680 p)	Contig132 (11041-12588 p)	Contig132 (10269-11021 p)	Contig132 (9698-10246 p)
SA-345.1	SA-347.1	SA-348.1	SA-351.1	SA-352.1	SA-353.1
SeqID 1730	SeqID 1731	SeqID 1732	SeqID 1733	SeqID 1734	SeqID 1735

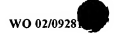


SeqID 1736	SA-356.1	Contig132 (8508-9530 p)	85	Identities = 257/339 (75%), Positives = 293/339 (85%), Gaps = 4/339 (1%) gb AAC16901.1  (AF016634) phosphoribosylformylglycinamide cyclo-ligase [Lactococcus lactis subsp. cremoris] Length = 340
SeqID 1737	SA-357.1	Contig132 (7026-8480 p)	. 8	Identities = 343/470 (72%), Positives = 406/470 (85%), Gaps = 6/470 (1%) pir  T51702 amidophosphoribosyltransferase (EC 2.4.2.14) [imported] - Lactococcus lactis gb AAD12627.1  (U64311) phosphoribosylpyrophosphate amidotransferase [Lactococcus lactis] Length = 506
SeqID 1738	SA-36.1	Contig137 (9548-10369 p)	No Hits found	
SeqID 1739	SA-361.1	Contig132 (4533-5015 m)	39	Identities = 39/137 (28%), Positives = 63/137 (45%), Gaps = 7/137 (5%) prf[ 2119294A YFW1 gene [Saccharomyces cerevisiae] Length = 605
SeqiD 1740	SA-363.1	Contig132 (3046-6792 p)	. 52	Identities = 198/746 (26%), Positives = 330/746 (43%), Gaps = 103/746 (13%) emb CAC12194.1  (AL445066) phosphoribosylformylglycinamidine synthase related protein [Thermoplasma acidophilum] Length = 759
SeqiD 1741	SA-364.2	Contig132 (2240-2944 p)	48	Identities = 183/235 (77%), Positives = 206/235 (86%) splQ07296 PUR7_STRPN PHOSPHORIBOSYLAMINOIMIDAZOLE- SUCCINOCARBOXAMIDE SYNTHASE (SAICAR SYNTHETASE) pir  A36941 phosphoribosylaminoimidazolesuccinocarboxamide synthase (EC 6.3.2.6) - Streptococcus pneumoniae gb AAA03540.1  (L15190) SAICAR synthetase [Streptococcus pneumoniae] gb AAA69512.1  (M36180) SAICAR synthetase [Streptococcus pneumoniae] Length = 235
SeqID 1742	SA-366.2	Contig139 (35877-38138 m)	85	Identities = 560/755 (74%), Positives = 648/755 (85%), Gaps = 12/755 (1%) gb AAD01782.1  (AF023421) ClpE [Lactococcus lactis] Length = 748



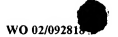
SeqID 1743	SA-367.1	Contig139 (35236-35691 m)	59	Identities = 31/101 (30%), Positives = 48/101 (46%), Gaps = 2/101 (1%) pir  A70315 AP4A hydrolase - Aquifex aeolicus gb AAC06510.1  (AE000676) AP4A hydrolase [Aquifex aeolicus] Length = 134
SeqID 1744	SA-368.1	Contig139 (34870-35172 m)	No Hits found	
SeqID 1745	SA-37.1	Contig137 (8461-9465 p)	37	Identities = 63/288 (21%), Positives = 129/288 (43%), Gaps = 27/288 (9%) ref[NP_053160.1  pXO2-05 [Bacillus anthracis] gb AAF13610.1 AF188935_7 (AF188935) pXO2-05 [Bacillus anthracis] Length = 282
SeqID 1746	SA-370.1	Contig139 (31961-34753 p)	87	Identities = 729/929 (78%), Positives = 821/929 (87%), Gaps = 1/929 (0%) splQ9ZHB3 SYI_STRPN ISOLEUCYL-TRNA SYNTHETASE (ISOLEUCINE-TRNA LIGASE) (ILERS) gb AAC95446.1  (AF068901) isoleucine-tRNA synthetase [Streptococcus pneumoniae] Length = 930
SeqID 1747	SA-371.1	. Contig139 (30906-31676 p)	89	Identities = 132/227 (58%), Positives = 179/227 (78%), Gaps = 2/227 (0%) gb AAC95445.1  (AF068901) cell division protein DivIVA [Streptococcus pneumoniae] Length = 262
SeqID 1748	SA-372.1	Contig139 (30108-30896 p)	09	Identities = 101/255 (39%), Positives = 162/255 (62%) gb[AAC95444.1] (AF068901) YImH [Streptococcus pneumoniae] Length = 261
SeqID 1749	SA-373.1	Contig139 (29852-30106 p)	29	Identities = 34/83 (40%), Positives = 54/83 (64%) emb CAA75619.1  (Y15422) hypothetical protein [Lactococcus lactis subsp. cremoris] Length = 91
SeqID 1750	SA-374.1	Contig139 (29244-29849 p)	25	Identities = 86/200 (43%), Positives = 120/200 (60%), Gaps = 25/200 (12%) gb AAC95442.1  (AF068901) YIMF [Streptococcus pneumoniae] Length = 179
SeqID 1751	SA-375.1	Contig139 (28558-29232 p)	76	Identities = 140/223 (62%), Positives = 177/223 (78%) gb AAC95441.1  (AF068901) YIME [Streptococcus pneumoniae] Length = 223
SeqID 1752	SA-376.1	Contig139 (27272-28552 p)	. 84	Identities = 327/426 (76%), Positives = 363/426 (84%), Gaps = 7/426 (1%) gb AAC95440.1  (AF068901) cell division protein FtsZ [Streptococcus pneumoniae] Length = 419

SeqID 1753	SA-377.1	Contig139 (25877-27250 p)	80	Identities = 293/458 (63%), Positives = 368/458 (79%), Gaps = 1/458 (0%) gblAAC95439.1 (AF068901) cell division protein FtsA [Streptococcus pneumoniae]
SeqID 1754	SA-379.1	Contig139 (24468-25604 p)	55	Identities = 123/396 (31%), Positives = 210/396 (52%), Gaps = 38/396 (9%) gb AAC95451.1  (AF068902) cell division protein DivIB [Streptococcus pneumoniae] Length = 399
SeqID 1755	SA-380.1	Contig139 (23388-24464 p)	08	Identities = 248/358 (69%), Positives = 293/358 (81%), Gaps = 1/358 (0%) dbj BAB16029.1  (AB030645) MurG [Streptococcus pyogenes] Length = 360
SeqID 1756	SA-381.1	Contig139 (22030-23385 p)	84	Identities = 340/449 (75%), Positives = 392/449 (86%) gb AAC95449.1  (AF068902) D-glutamic acid adding enzyme MurD [Streptococcus pneumoniae] Length = 450
SeqID 1757	SA-382.1	Contig139 (21655-21900 p)	No Hits found	
SeqID 1758	SA-384.1	Contig139 (19769-21610 p)		Identities = 457/609 (75%), Positives = 536/609 (87%), Gaps = 2/609 (0%) splO07631 TYPA_BACSU GTP-BINDING PROTEIN TYPA/BIPA HOMOLOG pir E69872 GTP-binding translation elongation factor homolog ylaG - Bacillus subtilis emb CAB09712.1  (297025) product highly similar to elongation factor EF-G [Bacillus subtilis] similar to GTP-binding elongation factor [Bacillus subtilis]
SeqID 1759	SA-385.1	Contig139 (19157-19537 p)	55	Identities = 51/124 (41%), Positives = 71/124 (57%), Gaps = 1/124 (0%) splP54510 YQHL_BACSU HYPOTHETICAL 14.6 KD PROTEIN IN GCVT-SPOIIIAA INTERGENIC REGION pir  C69959 glpE protein homolog yqhL - Bacillus subtilis dbj BAA12549.1  (D84432) YqhL [Bacillus subtilis] emb CAB14385.1  (299116) similar to hypothetical proteins [Bacillus subtilis]



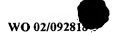
SeqID 1760	SA-387.1	Contig139 (18177-19145 p)	64	Identities = 145/315 (46%), Positives = 209/315 (66%), Gaps = 2/315 (0%) dbj BAB05144.1  (AP001512) glucose kinase [Bacillus halodurans] Length = 330
SeqID 1761	SA-388.1	Contig139 (17971-18180 p)	84	Identities = 53/67 (79%), Positives = 62/67 (92%) dbj BAA96473.1  (AB036428) hypothetical 8.3 kDa protein [Streptococcus mutans] Length = 68
SeqID 1762	SA-389.1	Contig139 (17421-17867 m)	57	Identities = 57/123 (46%), Positives = 85/123 (68%) dbj BAA96471.1  (AB036428) type IV prepilin peptidase homologue [Streptococcus mutans] Length = 218
SeqID 1763	SA-39.1	Contig137 (7426-8382 m)	No Hits found	
SeqID 1764	SA-390.1	Contig139 (16788-17420 p)	19	Identities = 95/202 (47%), Positives = 134/202 (66%)   dbj BAB05417.1  (AP001512) endonuclease III (DNA repair)   [Bacillus halodurans]   Length = 218   .
SeqID 1765	SA-391.1	Contig139 (15458-16675 p)	38	Identities = 88/384 (22%), Positives = 159/384 (40%), Gaps = 24/384 (6%) pir  A69832 probable acid-CoA ligase (EC 6.2.1) yhfT [similarity] - Bacillus subtilis emb CAA74543.1  (Y14084) hypothetical protein [Bacillus subtilis] emb CAB12876.1  (Z99109) similar to long-chain fatty-acid-CoA ligase [Bacillus subtilis] Length = 479
SeqID 1766	SA-392.1	Contig139 (14410-15453 p)	45	Identities = 96/340 (28%), Positives = 160/340 (46%), Gaps = 21/340 (6%) emb CAC11722.1  (AL445064) acetyl-CoA acetyltransferase related protein [Thermoplasma acidophilum] Length = 388
SeqID 1767	SA-393.1	Contig 139 (13834-14349 p)	No Hits found	
SeqID 1768	SA-394.1	Contig139 (12717-13709 m)	55	Identities = 132/316 (41%), Positives = 201/316 (62%), Gaps = 2/316 (0%) dbj BAB05467.1  (AP001513) biotin synthase [Bacillus halodurans] Length = 333

SeqID 1769	SA-395.1	Contig139 (12177-12716 m)	<b>8</b> 6.	Identities = 69/168 (41%), Positives = 105/168 (62%) pir  B69832 biotin biosynthesis homolog yhfU - Bacillus subtilis emb CAA74544.1  (Y14084) hypothetical protein [Bacillus subtilis] emb CAB12877.1  (299109) similar to biotin biosynthesis [Bacillus subtilis] Length = 186
SeqiD 1770	SA-396.1	Contig139 (11543-12109 p)	. 09	Identities = 102/191 (53%), Positives = 134/191 (69%), Gaps = 4/191 (2%) splQ02003 TRPG_LACLA ANTHRANILATE SYNTHASE COMPONENT II (GLUTAMINE AMIDO-TRANSFERASE) pir  S35125 anthranilate synthase (EC 4.1.3.27) beta chain - Lactococcus lactis subsp. lactis gb AAA25224.1  (M87483) anthranilate synthase beta subunit [Lactococcus lactis] Length = 198
SeqID 1771	SA-397.1	Contig139 (9673-11415 p)	89	identities = 280/582 (48%), Positives = 400/582 (68%), Gaps = 6/582 (1%) dbj BAB06054.1  (AP001515) ABC transporter (ATP-binding protein) [Bacillus halodurans] Length = 599
SeqID 1772	SA-398.1	Contig139 (7941-9686 p)	69	Identities = 284/575 (49%), Positives = 406/575 (70%), Gaps = 2/575 (0%) dbj BAB06055.1  (AP001515) ABC transporter (ATP-binding protein) [Bacillus halodurans] Length = 585
SeqID 1773	SA-399.1	Contig139 (7161-7745 p)	34	Identities = 46/154 (29%), Positives = 68/154 (43%), Gaps = 36/154 (23%) gb AAF36806.1 (AF155139) VanZF [Paenibacillus popilliae]
SeqID 1774	SA-4.1	Contig137 (39841-40488 p)	No Hits found	
SeqID 1775	SA-40.1	Contig137 (7185-7433 p)	No Hits found	
SeqID 1776	SA-400.1	Contig139 (6106-7230 p)	67	Identities = 206/349 (59%), Positives = 257/349 (73%), Gaps = 5/349 (1%) dbj BAB06225.1  (AP001515) unknown conserved protein [Bacillus halodurans]
SeqID 1777	SA-401.2	Contig139 (5495-6073 p)	29	Identities = 46/89 (51%), Positives = 62/89 (68%), Gaps = 1/89 (1%) pir F70023 hypothetical protein yutD - Bacillus subtilis emb CAB15221.1  (299120) yutD [Bacillus subtilis] Length = 102



, Gaps = in yunD - ar to = 462	, Gaps = 3acillus acillus subtilis]	!%) !NE PHATE 7-COA 01119)		Gaps = Inserved 19		os = 3/97 Bta 15	8%) LIGASE artate nzae rtate
Identities = 155/463 (33%), Positives = 262/463 (56%), Gaps = 33/463 (7%) pir  G70015 conserved hypothetical protein yunD - Bacillus subtilis emb CAB15227.1  (Z99120) similar to hypothetical proteins [Bacillus subtilis]	Identities = 125/344 (36%), Positives = 199/344 (57%), Gaps 21/344 (6%) pir[ A69875 hypothetical protein ylbL - Bacillus subtilis emb CAB11358.1 (298682) YlbL protein [Bacillus subtilis] emb CAB13378.1 (299111) ylbL [Bacillus subtilis] Length = 350	Identities = 56/149 (37%), Positives = 94/149 (62%) splP57643 COAD_BUCAI PHOSPHOPANTETHEINE ADENYLYLTRANSFERASE (PANTETHEINE-PHOSPHATE ADENYLYLTRANSFERASE) (PPAT) (DEPHOSPHO-COA PYROPHOSPHORYLASE) dbj BAB13272.1  (AP001119) lipopolysaccharide core biosynthesis protein kdtB [Buchnera sp. APS]		Identities = 80/180 (44%), Positives = 118/180 (65%), Gaps = 3/180 (1%) dbj BAB06309.1  (AP001516) unknown conserved protein [Bacillus halodurans]		Identities = 30/97 (30%), Positives = 51/97 (51%), Gaps = 3/97 (3%) gb AAD56628.1 AF165218_3 (AF165218) Bta [Streptococcus pneumoniae]	Identities = 267/328 (81%), Positives = 290/328 (88%) splP44338 ASNA_HAEIN ASPARTATEAMMONIA LIGASE (ASPARAGINE SYNTHETASE A) pir  H64077 aspartate-ammonia ligase (EC 6.3.1.1) - Haemophilus influenzae (strain Rd KW20) gb AAC22222.1  (U32738) aspartate-ammonia ligase (PC 6.3.1.1) - Haemophilus influenzae
. 28	55	25	No Hits found	46	No Hits found	31	98
Contig139 (4092-5426 p)	Contig139 (2944-4017 p)	Contig139 (2469-2954 p)	Contig139 (2149-2472 p)	Contig139 (1418-2170 p)	C. ntig137 (7096-7170 p)	Contig139 (1051-1419 m)	Contig139 (5-1006 p)
SA-402.2	SA-403.1	SA-405.1	SA-407.1	SA-409.1	SA-41.1	SA-410.1	SA-411.1
SeqID 1778	SeqID 1779	SeqID 1780	SeqID 1781	SeqID 1782	SeqID 1783	SeqID 1784	SeqID 1785

SeqID 1786	SA-412.2	Contig134 (36525-37172 m)	75	Identities = 125/212 (58%), Positives = 165/212 (76%)   gb AAF81675.1 AF232688_4 (AF232688) SioR (Streptococcus   mutans]
SeqID 1787	SA-413.1	Contig134 (35796-36485 p)	62	Identities = 100/229 (43%), Positives = 145/229 (62%) splP45113 MTN_HAEIN MTA/SAH NUCLEOSIDASE [INCLUDES: 5-METHYLTHIOADENOSINE NUCLEOSIDASE ; S-ADENOSYLHOMOCYSTEINE NUCLEOSIDASE] pir  C64169 pfs protein - Haemophilus influenzae (strain Rd KW20) gb AAC22869.1  (U32801) pfs protein (pfs) [Haemophilus influenzae Rd] Length = 229
SeqID 1788	SA-414.1	Contig134 (35517-35786 p)	No Hits found	
SeqID 1789	SA-415.1	Contig134 (34963-35517 p)	62	Identities = 92/181 (50%), Positives = 125/181 (68%), Gaps = 4/181 (2%) splP54570 YQKG_BACSU HYPOTHETICAL 21.0 KDA PROTEIN IN GLNQ-ANSR INTERGENIC REGION pir  A69967 conserved hypothetical protein yqkG - Bacillus subtilis dbj BA412639.1  (D84432) YqkG [Bacillus subtilis] emb CAB14293.1  (299116) similar to hypothetical proteins [Bacillus subtilis]
SeqID 1790	SA-416.1	Contig134 (33563-34942 p)	89	Identities = 238/453 (52%), Positives = 322/453 (70%), Gaps = 1/453 (0%) dbj BAB03784.1  (AP001507) UDP-N-acetylglucosamine pyrophosphorylase [Bacillus halodurans] Length = 455
SeqID 1791	SA-417.1	Contig134 (32906-33319 p)	28	Identities = 46/144 (31%), Positives = 83/144 (56%), Gaps = 10/144 (6%) dbj BAB04569.1  (AP001510) unknown conserved protein in others [Bacillus halodurans] Length = 148
SeqID 1792	SA-418.1	Contig134 (32516-32893 p)	48	Identities = 41/127 (32%), Positives = 63/127 (49%), Gaps = 11/127 (8%) pir  E69798 conserved hypothetical protein yetH - Bacillus subtilis emb CAB12535.1 (Z99107) similar to hypothetical proteins [Bacillus subtilis] Length = 120
SeqID 1793	SA-419.1	Contig134 (31471-32427 p)	. 09	Identities = 133/320 (41%), Positives = 193/320 (59%), Gaps = 4/320 (1%) dbj BAB06422.1  (AP001516) unknown conserved protein [Bacillus halodurans]

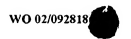


	Identities = 27/90 (30%), Positives = 49/90 (54%), Gaps = 7/90 (7%) pirl T35570 hypothetical protein SC6G4.19c SC6G4.19c Streptomyces coelicolor emb CAA20397.1  (AL031317) SC6G4.19c, unknown, len: 190 aa; contains Pro-Ser- r ich domain at N-terminus [Streptomyces coelicolor A3(2)] Length = 190			Identities = 82/239 (34%), Positives = 125/239 (51%), Gaps = 15/239 (6%) pir[ T44434 3-oxoacyl-[acyl-carrier-protein] reductase (EC 1.1.1.100) [imported] - Moritella marina dbj BAA85256.1  (AB021978) 3-oxoacyl-[acyl carrier protein] reductase homolog [Moritella marina] Length = 244			Identities = 35/153 (22%), Positives = 64/153 (40%), Gaps = 5/153 (3%) pir  T35964 hypothetical protein SC9C7.13c - Streptomyces coelicolor emb CAA22725.1  (AL035161) hypothetical protein SC9C7.13c [Streptomyces coelicolor A3(2)] Length = 179	Identities = 30/138 (21%), Positives = 68/138 (48%) dbj BAB05950.1  (AP001514) unknown conserved protein in others [Bacillus   halodurans] Length = 482	Identities = 126/320 (39%), Positives = 180/320 (55%), Gaps = 18/320 (5%) dbj BAB05950.1  (AP001514) unknown conserved protein in others [Bacillus halodurans]
No Hits found		No Hits found	No Hits found No Hits found		No Hits found	No Hits found	33	44	55
Contig137 (6439-6849 p)	Contig134 (31220-31474 p)	Contig134 (31045-31359 m)	Contig134 (31017-31259 p) Contig134 (30787-31020 p)	Contig134 (30080-30778 p)	Contig134 (29851-30087 p)	Contig134 (29501-29761 p)	Contig134 (28815-29315 p)	Contig134 (28120-28575 p)	. Contig134 (27217-28119 p)
SA-42.1	SA-420.1	SA-421.1	SA-422.1 SA-423.1	SA-424.1	SA-425.1	SA-426.1	SA-427.1	SA-428.1	SA-429.1
SeqID 1794	SeqiD 1795	SeqID 1796	SeqID 1797 SeqID 1798	SeqID 1799	SeqID 1800	SeqID 1801	SeqID 1802	SeqID 1803	SeqID 1804

Identities = 199/697 (28%), Positives = 322/697 (45%), Gaps = 58/697 (8%) dbjjBAB05949.1  (AP001514) unknown [Bacillus halodurans] Length = 1091	Identities = 71/359 (19%), Positives = 148/359 (40%), Gaps = 49/359 (13%) splQ45535 COTH_BACSU INNER SPORE COAT PROTEIN H pir  E69605 spore coat protein (inner) cotH - Bacillus subtilis emb CAB07793.1  (293767) ywrH [Bacillus subtilis] emb CAB15623.1  (299122) spore coat protein (inner) [Bacillus subtilis]			Identities = 162/436 (37%), Positives = 248/436 (56%), Gaps = 13/436 (2%) pir[ C69596 branched-chain amino acid transporter braB - Bacillus subtilis gb AAC00400.1  (AF008220) branch-chain amino acid transporter [Bacillus subtilis] emb CAB14938.1  (Z99119) branched-chain amino acid transporter [Bacillus subtilis] Length = 445	Identities = 395/667 (59%), Positives = 501/667 (74%), Gaps = 12/667 (1%) splP37465 SYM_BACSU METHIONYL-TRNA SYNTHETASE (METHIONINETRNA LIGASE) (METRS) pir  S66067 methioninetRNA ligase (EC 6.1.1.10) metS - Bacillus subtilis dbj BAA05273.1  (D26185) methionyl-tRNA synthetase [Bacillus subtilis] emb CAB11814.1  (299104) methionyl-tRNA synthetase [Bacillus subtilis]	Identities = 163/282 (57%), Positives = 205/282 (71%), Gaps = 1/282 (0%) splP45134 TEHB_HAEIN TELLURITE RESISTANCE PROTEIN TEHB HOMOLOG pir  H64113 hemagglutinin hag1 homolog - Haemophilus influenzae (strain Rd KW20) gb AAC22923.1  (U32807) tellurite resistance protein (tehB) [Haemophilus influenzae Rd] Length = 286
44	. 29	No Hits found	No Hits found	55	72	69
Contig134 (24924-27023 p)	Contig134 (23373-24857 p)		$\overline{}$	Contig134 (20288-21631 p)	Contig134 (18101-20098 p)	Contig134 (17083-17958 m)
SA-430.1	SA-431.1	SA-432.1	SA-433.1	SA-434.2	SA-435.1	SA-436.1
SeqID 1805	SeqID 1806	SeqID 1807	SeqID 1808	SeqID 1809	SeqID 1810	SeqID 1811



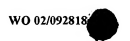
Contig134 (16203-16925 p)
Contig134 (14677-15894 m)
Contig134 (13851-14636 m)
Contig137 (5997-6446 p) No Hits found
Contig134 (12947-13774 m)
Contig134 (12550-12906 p)
Contig134 (12072-12548 p)



SeqID 1819	SA-445.1	Contig134 (10835-12016 p)	42	Identities = 102/313 (32%). Positives = 168/313 (53%), Gaps = 21/313 (6%) splO58424 SERA_METJA D-3-PHOSPHOGLYCERATE DEHYDROGENASE (PGDH) pir[ A64427 phosphoglycerate dehydrogenase (EC 1.1.1.95) - Methanococcus jannaschii gb AAB99020.1  (U67544) phosphoglycerate dehydrogenase (serA) [Methanococcus jannaschii] Length = 524
SeqID 1820	SA-446.1	Contig134 (10213-10773 p)	41	Identities = 45/170 (26%), Positives = 78/170 (45%), Gaps = 13/170 (7%) pir  G81269 probable acetyltransferase Cj1715 [imported] - Campylobacter jejuni (strain NCTC 11168) emb  CAB73701.1  (AL139079) putative acetyltransferase [Campylobacter jejuni]
SeqID 1821	SA-447.1	Contig134 (9065-10156 p)		Identities = 169/363 (46%), Positives = 252/363 (68%), Gaps = 8/363 (2%) gbJAAF13453.1JAF204962_1 (AF204962) phosphoserine aminotransferase [Bacillus alcalophilus] Length = 361
SeqID 1822	SA-448.1	Contig134 (8297-8932 m)	. 54	Identities = 80/203 (39%), Positives = 116/203 (56%), Gaps = 7/203 (3%) ref[XP_005810.1  CGI-32 protein [Homo sapiens] Length = 307
SeqID 1823	SA-45.1	Contig137 (5361-5807 p)	No Hits found	
SeqID 1824	SA-450.1	Contig134 (7155-8027 p)	65	Identities = 141/287 (49%), Positives = 190/287 (66%), Gaps = 2/287 (0%) dbj BAB03768.1  (AP001507) unknown conserved protein [Bacillus halodurans] Length = 289
SeqiD 1825	SA-451.1	Contig134 (6832-7158 p)	56	Identities = 45/116 (38%), Positives = 62/116 (52%), Gaps = 8/116 (6%) dbj BAB03765.1  (AP001507) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 116
SeqID 1826	SA-452.1	Contig134 (5938-6801 p)	. 14	Identities = 82/219 (37%), Positives = 124/219 (56%), Gaps = 3/219 (1%) dbj BAB03763.1  (AP001507) DNA polymerase III delta subunit [Bacillus halodurans] Length = 328



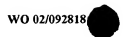
		<del>,</del>		<del></del>
Identities = 112/210 (53%), Positives = 148/210 (70%), Gaps = 1/210 (0%) dbj BAB03761.1  (AP001507) thymidylate kinase [Bacillus halodurans] Length = 210	Identities = 73/166 (43%), Positives = 116/166 (68%), Gaps = 2/166 (1%) pir  H72290 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD36216.1 AE001771_9 (AE001771) conserved hypothetical protein [Thermotoga maritima] Length = 215	Identities = 135/233 (57%), Positives = 180/233 (76%) pir  T35757 probable branched chain amino acid transport ATP- binding protein - Streptomyces coelicolor emb CAB52068.1  (AL109732) putative branched chain amino acid transport ATP-binding protein [Streptomyces coelicolor A3(2)] Length = 238	Identities = 136/273 (49%), Positives = 190/273 (68%), Gaps = 21/273 (7%) pir] F72290 branched chain amino acid ABC transporter, ATP-binding protein - Thermotoga maritima (strain MSB8) gb AAD36214.1 AE001771_7 (AE001771) branched chain amino acid ABC transporter, ATP-binding protein [Thermotoga maritima]	Identities = 121/343 (35%), Positives = 195/343 (56%), Gaps = 36/343 (10%) pir  E72290 branched chain amino acid ABC transporter, permease protein - Thermotoga maritima (strain MSB8) gb AAD36213.1 AE001771_6 (AE001771) branched chain amino acid ABC transporter, permease protein [Thermotoga maritima] Length = 359
59	51	75	74	09
Contig134 (5283-5918 p)	Contig134 (4535-5194 p)	Contig134 (3806-4516 p)	Contig134 (3042-3806 p)	Contig134 (2088-3041 p)
SA-453.1	SA-454.1	SA-455.1	SA-456.1	SA-457.1
SeqID 1827	SeqID 1828	SeqID 1829	SeqID 1830	SeqID 1831



Identities = 14U/295 (47%), Positives = 200/295 (67%), Gaps = 7/295 (2%) pir  D72290 branched chain amino acid ABC transporter, permease protein - Thermotoga maritima (strain MSB8) gb AAD36212.1 AE001771_5 (AE001771) branched chain amino acid ABC transporter, permease protein [Thermotoga maritima] Length = 299	No Hits found	Identities = 114/360 (31%), Positives = 182/360 (49%), Gaps = 27/360 (7%) pir[]C72290 branched chain amino acid ABC transporter, periplasmic amino acid-binding protein - Thermotoga maritima (strain MSB8) gb]AAD36211.1]AE001771_4 (AE001771) branched chain amino acid ABC transporter, periplasmic amino acid-binding protein [Thermotoga maritima] Length = 370	Identities = 30/102 (29%), Positives = 60/102 (58%)   gb AAD05186.1  (AF110185) unknown [Burkholderia   pseudomallei] Length = 163	Identities = 224/324 (69%), Positives = 276/324 (85%), Gaps = 1/324 (0%) gb AAF98271.1 AF197933_1 (AF197933) betaketoacyl-ACP synthase III [Streptococcus pneumoniae]  Length = 324	Identities = 64/74 (86%), Positives = 67/74 (90%)   20 AAF98272.1 AF197933_2 (AF197933) acyl carrier protein   Streptococcus pneumoniae	Identities = 257/318 (80%), Positives = 277/318 (86%), Gaps = 1/318 (0%) gb AAF98273.1 AF197933_3 (AF197933) trans-2-enoyl-ACP reductase II [Streptococcus pneumoniae]  Length = 324
	No Hits				•	
Contig134 (1216-2085 p)	Contig137 (4933-5361 p)	Contig134 (1-1110 p)	Contig135 (65101-65484 m)	Contig135 (64130-65101 m)	Contig135 (63848-64072 m)	Contig135 (62734-63693 m)
SA-458.2	SA-46.1	SA-460.2	SA-462.1	SA-463.1	SA-464.1	SA-465.1
SeqID 1832	SeqID 1833	SeqID 1834	SeqID 1835	SeqID 1836	SeqID 1837	SeqiD 1838



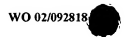
SeqID 1839	SA-466.1	Contig135 (61788-62714 m)	77	Identities = 204/306 (66%), Positives = 242/306 (78%), Gaps = 1/306 (0%) gb AAF98274.1 AF197933_4 (AF197933) malonyl-CoA:ACP transacylase; MCAT [Streptococcus pneumoniae] Length = 306
SeqID 1840	SA-467.1	Contig135 (61045-61779 m)	84	Identities = 184/243 (75%), Positives = 212/243 (86%)   gb AAF98275.1 AF197933_5 (AF197933) beta-ketoacyl-ACP   reductase [Streptococcus pneumoniae]
SeqID 1841	SA-468.1	Contig135 (59797-61029 m)	06	Identities = 340/410 (82%), Positives = 375/410 (90%)   gb AAF98276.1 AF197933_6 (AF197933) beta-ketoacyl-ACP   synthase II [Streptococcus pneumoniae] Length = 411
SeqID 1842	SA-469.1	Contig135 (59295-59795 m)	58	Identities = 103/169 (60%), Positives = 127/169 (74%), Gaps = 11/169 (6%) gb AAF98277.1 AF197933_7 (AF197933) biotin carboxyl carrier protein [Streptococcus pneumoniae]
SeqID 1843	SA-47.1	Contig137 (2969-4462 p)	73	Identities = 278/497 (55%), Positives = 367/497 (72%), Gaps = 1/497 (0%) pir JH0206 hypothetical 57.4K protein - Enterococcus faecalis plasmid pAM-beta-1 gb AAB94754.1  (AF039139) replication protein E [Cloning vector plL252] gb AAB96788.1  (AF041239) replication protein E [Cloning vector plL253] gb AAC38603.1  (AF007787) RepE [Enterococcus faecalis] Length = 496
SeqID 1844	SA-470.1	Contig135 (58876-59298 m)	91	Identities = 130/140 (92%), Positives = 135/140 (95%) gb AAF98278.1 AF197933_8 (AF197933) beta-hydroxyacyl-ACP dehydratase [Streptococcus pneumoniae] Length = 140
SeqID 1845	SA-471.1	Contig135 (57468-58838 m)	85	Identities = 361/451 (80%), Positives = 405/451 (89%) gb AAF98279.1 AF197933_9 (AF197933) acetyl-CoA carboxylase biotin carboxylase subunit [Streptococcus pneumoniae] Length = 455



SeqID 1846	SA-472.1	Contig135 (56584-57459 m)	83	Identities = 221/285 (77%), Positives = 248/285 (86%), Gaps = 1/285 (0%) gb AAF98280.1 AF197933_10 (AF197933) acetyl-CoA carboxylase beta subunit [Streptococcus pneumoniae] Length = 288
SeqID 1847	SA-473.1	Contig135 (55818-56591 m)	83	Identities = 186/254 (73%), Positives = 222/254 (87%) gb AAF98281.1 AF197933_11 (AF197933) acetyl-CoA carboxylase alpha subunit [Streptococcus pneumoniae] Length = 255
SeqID 1848	SA-474.1	Contig135 (54716-55264 m)	45	Identities = 76/142 (53%), Positives = 99/142 (69%) dbj BAA88824.1  (AB016077) sakacin A production response regulator [Streptococcus mutans]
SeqID 1849	SA-475.1	Contig135 (53393-54670 p)	74	Identities = 262/425 (61%), Positives = 322/425 (75%), Gaps = 1/425 (0%) splP37464 SYS_BACSU SERYL-TRNA SYNTHETASE (SERINETRNA LIGASE) (SERRS) pirl S66043 serinetRNA ligase (EC 6.1.1.11) - Bacillus subtilis dbj BAA05249.1  (D26185) seryl-tRNA synthetase [Bacillus subtilis] emb CAB11789.1  (Z99104) seryl-tRNA synthetase [Bacillus subtilis] Length = 425
SeqID 1850	SA-477.1	Contig135 (52035-53102 m)	40	Identities = 72/330 (21%), Positives = 143/330 (42%), Gaps = 32/330 (9%) emb CAA07406.1  (AJ006986) transmembrane protein [Streptococcus pneumoniae] Length = 332
SeqID 1851	SA-478.1	Contig135 (51635-51997 p)	75	Identitles = 88/112 (78%), Positives = 96/112 (85%) gbjAAD46488.1 AF130465_4 (AF130465) unknown [Streptococcus salivarius] Length = 124
SeqID 1852	SA-48.1	Contig137 (2779-2871 m)	No Hits found	
SeqID 1853	SA-480.1	Contig135 (50605-51516 p)	06	Identities = 247/303 (81%), Positives = 276/303 (90%) gb AAD46487.1 AF130465_3 (AF130465) mannose-specific phosphotransferase system component IID [Streptococcus salivarius] Length = 303
SeqID 1854	SA-481.1	Contig135 (49778-50590 p)	84	Identities = 209/271 (77%), Positives = 233/271 (85%), Gaps = 1/271 (0%) gb AAD46486.1 AF130465_2 (AF130465) mannosespecific phosphotransferase system component IIC [Streptococcus salivarius]



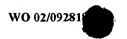
	SA-483.1	Contig135 (48735-49745 p)	68	Identities = 287/336 (85%), Positives = 306/336 (90%), Gaps = 6/336 (1%) gblAAD46485.1IAF130465_1 (AF130465) mannosespecific phosphotransferase system component IIAB (Streptococcus salivarius) Length = 330
	SA-484.1	Contig135 (47620-48432 p)	20	Identities = 89/267 (33%), Positives = 139/267 (51%), Gaps = 3/267 (1%) dbj BAB06625.1  (AP001517) unknown conserved protein [Bacillus halodurans] Length = 269
	SA-485.1	Contig135 (46947-47531 m)	No Hits found	
	SA-486.1	Contig135 (46246-46773 m)	4	Identities = 52/189 (27%), Positives = 92/189 (48%), Gaps = 12/189 (6%) pir] C71375 conserved hypothetical integral membrane protein TF0033 - syphilis spirochete gb AAC65028.1  (AE001188) conserved hypothetical integral membrane protein [Treponema pallidum] Length = 203
	SA-487.1	Contig135 (45206-45931 p)	No Hits found	
	SA-488.1	Contig135 (44722-46143 m)	58	Identities = 193/471 (40%), Positives = 286/471 (59%), Gaps = 42/471 (8%) pir  B82096 conserved hypothetical protein VC2278 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF95422.1  (AE004299) conserved hypothetical protein [Vibrio cholerae] Length = 430
	SA-489.1	Contig135 (44130-44573 m)	53	Identities = 68/149 (45%), Positives = 101/149 (67%), Gaps = 1/149 (0%) dbj BAB04264.1  (AP001508) unknown conserved protein [Bacillus halodurans]
	SA-49.2	Contig137 (2519-2611 p)	No Hits found	
	SA-490.1	Contig135 (43616-44137 m)	6	Identities = 56/163 (34%), Positives = 94/163 (57%), Gaps = 1/163 (0%) pir] D72360 conserved hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD35662.1 AE001732_7 (AE001732) conserved hypothetical protein [Thermotoga maritima] Length = 179



SeqID 1864	SA-491.1	Contig135 (42300-43607 m)	. 40	Identities = 108/324 (33%), Positives = 178/324 (54%), Gaps = 33/324 (10%) sp[Q02115]LYTR_BACSU MEMBRANE-BOUND PROTEIN LYTR pir][A47679 by divergon expression attenuator LytR - Bacillus subtilis gb[AAA22578.1] (M87645) membrane bound protein [Bacillus subtilis] emb[CAB15582.1] (299122) membrane-bound protein [Bacillus subtilis] Length = 306
SeqID 1865	SA-493.1	Contig135 (41940-42236 p)	No Hits found	
SeqID 1866	SA-494.1	Contig135 (41524-41943 p)	65	Identities = 74/137 (54%), Positives = 95/137 (69%), Gaps = 2/137 (1%) dbj BAB04908.1  (AP001511) Hit-like protein involved in cell-cycle regulation [Bacillus halodurans] Length = 142
SeqID 1867	SA-495.1	Contig 135 (40685-41272 m)	No Hits found	
SeqID 1868	SA-497.1	Contig135 (38248-40410 p)	99	Identities = 381/652 (58%), Positives = 485/652 (73%), Gaps = 15/652 (2%) spl034580 PCRA_BACSU ATP-DEPENDENT DNA HELICASE PCRA pir  E69794 ATP-dependent DNA helicase homolog yerF - Bacillus subtilis emb CAA75552.1 (Y15254) PcrA protein [Bacillus subtilis] emb CAB12481.1 (299107) similar to ATP-dependent DNA helicase [Bacillus subtilis]
SeqID 1869	SA-498.1	Contig135 (37777-38163 p)	53	/ Identitles = 61/87 (70%), Positives = 71/87 (81%) gb AAA88579.1  (M14339) unknown [Streptococcus pneumoniae] Length = 93
SeqID 1870	SA-499.1	Contig135 (36427-37644 p)	62	Identities = 208/423 (49%), Positives = 293/423 (69%), Gaps = 11/423 (2%) sp P41006 PYRP_BACCL URACIL PERMEASE (URACIL TRANSPORTER) pir  538893 uracil transport protein - Bacillus caldolyticus emb CAA53697.1  (X76083) uracil permease [Bacillus caldolyticus]
SeqID 1871	SA-5.1	Contig137 (39530-39841 p)	No Hits found	
SeqID 1872	SA-50.1	Contig137 (2617-2736 p)	No Hits found	



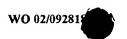
	No Hits found	COITING 135 (29390-29749 III)	1.700-20	ו מפטי היושפי
	No Hits found	Contig135 (30121-30657 m)	SA-506.1	SeqID 1879
Identities = 44/194 (22%), Positives = 90/194 (45%), Gaps = 13/194 (6%) emb CAA76857.1  (Y17797) hypothetical protein [Enterococcus faecalis] Length = 247	42	Contig135 (31153-31782 m)	SA-504.1	SeqID 1877
Identities = 80/226 (35%), Positives = 136/226 (59%), Gaps = 1/226 (0%) pir  D69983 conserved hypothetical protein ysbB - Bacillus subtilis emb CAA99613.1  (Z75208) hypothetical protein [Bacillus subtilis] emb CAB14850.1  (Z99118) similar to hypothetical proteins [Bacillus subtilis]	53	Contig135 (31850-32545 m)	SA-503.1	SeqID 1876
Identities = 34/110 (30%), Positives = 67/110 (60%), Gaps = 1/110 (0%) pir  A71191 hypothetical protein PH1801 - Pyrococcus horikoshii dbj BAA30920.1  (AP000007) 109aa long hypothetical protein [Pyrococcus horikoshii] Length = 109	49	Contig135 (32526-32918 m)	SA-502.1	SeqID 1875
Identities = 118/282 (41%), Positives = 181/282 (63%) splP46348 YEAB_BACSU HYPOTHETICAL 31.8 KD PROTEIN IN GABP-GUAA INTERGENIC REGION (ORFX) pir  B69791 cation efflux system membrane protein homolog yeaB - Bacillus subtilis; gb AAB62307.1  (U51115) YeaB [Bacillus subtilis] emb CAB12451.1  (Z99107) alternate gene name: ydxT~similar to cation efflux system membrane protein [Bacillus subtilis]	64	Contig135 (33077-34312 m)	SA-501.1	SeqID 1874
Identities = 256/443 (57%), Positives = 333/443 (74%), Gaps = 4/443 (0%) sp P44917 Y883_HAEIN HYPOTHETICAL PROTEIN HI0883 pirl H64099 probable amino acid transport protein H10883, sodium-dependent - Haemophilus influenzae (strain Rd KW20) gb AAC22541.1  (U32770) amino acid carrier protein, putative [Haemophilus influenzae Rd]  Length = 456	69	Contig135 (34376-35722 m)	SA-500.1	SeqID 1873



	Contig135 (28209-28997 m)	53	Identities = 124/246 (50%), Positives = 161/246 (65%), Gaps = 22/246 (8%) pir] C28551 hypothetical protein 3 - Streptococcus mutans (strain GS-5) (fragment) gb AAA88585.1  (M18954) unknown protein [Streptococcus mutans] Length = 228
Co	Contig135 (73667-74551 p)	99	Identities = 150/285 (52%), Positives = 198/285 (68%), Gaps = 5/285 (1%) gb AAK04462.1 AE006273_8 (AE006273) pseudouridine synthase [Lactococcus lactis subsp. Length = 293
3	Contig137 (2184-2513 p)	No Hits found	
Conti	Contig135 (72831-73670 p)	61	Identities = 120/267 (44%), Positives = 174/267 (64%), Gaps = 3/267 (1%) splO31612 YJBN_BACSU HYPOTHETICAL 30.0 KDA PROTEIN IN MECA-TENA INTERGENIC REGION pir  F69844 conserved hypothetical protein yjbN - Bacillus subtilis emb CAB13018.1  (299110) similar to hypothetical proteins [Bacillus subtilis]
Contig135	g135 (72188-72859 p)	69	dentities = 115/208 (55%), Positives = 159/208 (76%), Gaps = 3/208 (1%) dbj BAB06568.1  (AP001516) GTP pyrophosphokinase [Bacillus halodurans] Length = 211
Contig 135	135 (71506-72078 m)	46	identities = 63/184 (34%), Positives = 99/184 (53%), Gaps = 10/184 (5%) pir  C69844 hypothetical protein yjbK - Bacillus subtilis emb  CAB13015.1  (Z99110) yjbK   Bacillus subtilis  Length = 190



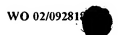
Identities = 166/319 (52%). Positives = 231/319 (72%). Gaps = 4/319 (1%) splP 14193 RPRS_BACSU RIBOSE-PHOSPHATE PYROPHOSPHOKINASE (PHOSPHORIBOSYL PYROPHOSPHATE SYNTHETASE) prijkliBSRS ribose-phosphate SYNTHETASE Of Allosteric Inhibition And Activation. pdbj1DKUJB Chain B, Crystal Structures Of Bacillus Subtilis Phosphoribosylpyrophosphate Synthetase: Molecular Basis Of Allosteric Inhibition And Activation. pdbj1DKUJB Chain B, Crystal Structures Of Bacillus Subtilis Phosphoribosylpyrophosphate Synthetase: Molecular Basis Of Allosteric Inhibition And Activation. pdbj1DKUJB Chain B, Crystal Structures Of Bacillus Subtilis Phosphoribosylpyrophosphate Synthetase: Molecular Basis Of Allosteric Inhibition And Activation. pdbj1DKUJB Chain B, Crystal Structures Of Bacillus Subtilis Phosphoribosylpyrophosphate Synthetase: Molecular Basis Of Allosteric Inhibition And Activation. pdbj1DKUJB CA94523.1   (X16518) PRPP Synthetase (AA	Identitites = 63/118 (53%), Positives = 81/118 (68%), Gaps = 1/118 (6%) gb AAG01802.1 AF276772_1 (AF276772) cysteine desulfurase NifS [Methanosarcina thermophila] Length = 404	Contig135 (69318-69647 m) No Hits found	Identities = 223/448 (49%), Positives = 313/448 (69%)  splP94417 AK3_BACSU PROBABLE ASPARTOKINASE (ASPARTATE KINASE) pir  A69763 homoserine dehydrogenase homolog yclM - Bacillus subtilis dbj BAA09011.1  (D50453) homologue of aspartokinase 2 alpha and beta subunits LysC of B. subtilis [Bacillus subtilis] emb CAB12187.1  (Z99106) similar to homoserine dehydrogenase [Bacillus subtilis]
SA-513.1	SA-514.1	SA-515.1	SA-517.1
SeqID 1887	SeqID_1888	SeqID 1889	SeqID 1890



SeqiD 1891	SA-518.1	Contig135 (66559-67209 m)	42	Identities = 59/190 (31%), Positives = 93/190 (48%), Gaps = 10/190 (5%) pir  A82498 CbbY-family-protein-VGA0102 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF96016.1  (AE004353) CbbY family protein [Vibrio cholerae] - Length = 2.19
SeqID 1892	SA-519.3	Contig135 (65631-66422 m)	64	Identities = 139/248 (56%), Positives = 185/248 (74%) dbj BAB16033.1  (AB030809) Pseudomonas putida enoyl-CoA hydratase II homologue [Streptococcus pyogenes] Length = 248
SeqID 1893	SA-52.1	Contig137 (1679-2173 p)	No Hits found	
SeqID 1894	SA-520.2	Contig116 (395-688 m)	42	Identities = 24/86 (27%), Positives = 43/86 (49%), Gaps = 6/86 (6%) gb AAD33114.1 AF094574_4 (AF094574) negative regulator of translation [Haemophilus influenzae]
SeqID 1895	SA-523.2	Contig116 (1354-4734 m)	73	dentities = 767/1103 (69%), Positives = 836/1103 (75%), Gaps = 58/1103 (5%) gb AAD39085.1 AF091393_1 (AF091393) surface protein R28 [Streptococcus pyogenes] Length = 1260
SeqID 1896	SA-524.1	Contig116 (4977-6170 p)	45	Identities = 187/187 (100%), Positives = 187/187 (100%) gb AAG09971.1 AF248037_6 (AF248037) unknown [Streptococcus agalactiae] Length = 189
SeqID 1897	SA-526.1	Contig116 (6365-6889 m)	<i>2</i> 6	Identitites = 174/174 (100%), Positives = 174/174 (100%) gb[AAG09969.1 AF248037_4 (AF248037) unknown [Streptococcus agalactiae] Length = 174
SeqID 1898	SA-527.2	Contig116 (7021-7599 p)	06	ss = 1{ 18037_ s agala
SeqID 1899	SA-528.2	Contig116 (7971-8354 p)	92	Identities = 125/128 (97%), Positives = 128/128 (99%)   gb AAG09967.1 AF248037_2 (AF248037) aldose reductase   [Streptococcus agalactiae]
SeqID 1900	SA-529.1	Contig116 (8355-8825 p)	96	Identities = 152/156 (97%), Positives = 152/156 (97%), Gaps = 4/156 (2%) gb AAG09967.1 AF248037_2 (AF248037) aldose reductase [Streptococcus agalactiae] Length = 280



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	dehydrogenase [Streptococcus agalactiae] Length = 96	Identities = 114/231 (49%), Positives = 161/231 (69%), Gaps = 6/231 (2%) gb AAG20655.1  (AE005134) alcohol dehydrogenase; Adh2 [Halobacterium sp. NRC-1] Length = 347	Identities = 53/123 (43%), Positives = 84/123 (68%) pir  B69970 transcription regulator MerR family homolog yraB - Bacillus subtilis emb CAA63468.1  (X92868) mercuric resistance operon regulatory protein [Bacillus subtilis] emb CAB14642.1  (Z99117) similar to transcriptional regulator (MerR family) [Bacillus subtilis] Length = 140	Identities = 55/131 (41%), Positives = 73/131 (54%), Gaps = 8/131 (6%) pir] B72308 hypothetical protein - Thermotoga maritima (strain MSB8) gb AAD36075.1 AE001762_2 (AE001762) hypothetical protein [Thermotoga maritima] Length = 135	Identities = 33/93 (35%), Positives = 59/93 (62%), Gaps = 1/93 (1%) pir  T29425 4-carboxymuconolactone decarboxylase homolog - Streptomyces coelicolor emb CAA20070.1  (AL031155) 3-oxoadipate enol-lactone hydrolase/4-carboxymuconolactone decarboxylase [Streptomyces coelicolor A3(2)] Length = 449	
No Hits found	28			54	48	No Hits found
Contig137 (1454-1642 p)	Contig116 (8935-9390 p)	Contig116 (9315-9971 p)	Contig116 (9987-10376 p)	Contig116 (10386-10781 p)	Contig116 (10805-11122 p)	Contig 116 (11209-11493 p)
SA-53.1	SA-530.1	SA-531.1	SA-532.1	SA-533.1	SA-534.1	SA-535.1
SeqID 1901	SeqID 1902	SeqID 1903	SeqID 1904	SeqID 1905	SeqID 1906	SeqID 1907



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90 Kg 69 69 69 69 69 69 69 69 69 69 69 69 69	Identities = 42/130 (32%), Positives = 71/130 (54%), Gaps = 3/130 (2%) pir  G69153 conserved hypothetical protein MTH413 - Methanobacterium thermoautotrophicum (strain Delta H) gb AAB84919.1  (AE000825) conserved protein [Methanobacterium thermoautotrophicum]	Identities = 141/566 (24%), Positives = 242/566 (41%), Gaps = 52/566 (9%) pir  T31094 surface antigen BspA - Bacteroides forsythus gb AAC82625.1  (AF054892) surface antigen BspA   [Bacteroides forsythus] Length = 1081	Identities = 102/413 (24%), Positives = 189/413 (45%), Gaps = 76/413 (18%) splP55340 ECSB_BACSU PROTEIN ECSB pir  G69619 ABC transporter (membrane protein) ecsB - Bacillus subtilis emb CAA61075.1  (X87807) hypothetical EcsB protein [Bacillus subtilis] emb CAA74408.1  (Y14077) Hypothetical protein [Bacillus subtilis] emb CAB12845.1  (299109) ABC transporter (membrane protein) [Bacillus subtilis] Length = 408	Identities = 475/727 (65%), Positives = 585/727 (80%), Gaps = 19/727 (2%) splP78027 RIR1_MYCPN RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE ALPHA CHAIN (RIBONUCLEOTIDE REDUCTASE) pir.  S73838 ribonucleoside-diphosphate reductase (EC 1.17.4.1) large chain nrdE - Mycoplasma pneumoniae (strain ATCC 29342) gb AAB96160.1  (AE000050) ribonucleoside-diphosphate reductase alpha chain-MPN324(new), 513(Himmelreich et al., 1996) [Mycoplasma pneumoniae] Length = 721
42	52	22	53.	. 49
Contig116 (11732-12223 p)	Contig116 (12267-12656 m)	Contig116 (12669-15836 m)	Contig137 (321-1361 p)	Contig116 (15899-18067 m)
SA-536.1	SA-537.1	SA-538.1	SA-54.1	SA-540.1
SeqID 1908	SeqiD 1909	SeqID 1910	SeqiD 1911	SeqiD 1912



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Identities = 76/127 (59%), Positives = 105/127 (81%), Gaps = 1/127 (0%) sp P47472 NRDI_MYCGE-NRDI-PROTEIN pir  D64225 hypothetical protein MG230 - Mycoplasma genitalium gb AAC71451.1  (U39702) nrdl protein (nrdl) [Mycoplasma genitalium]	Identities = 259/335 (77%), Positives = 299/335 (88%) sp P75461 RIR2_MYCPN RIBONUCLEOSIDE-DIPHOSPHATE REDUCTASE BETA CHAIN (RIBONUCLEOTIDE REDUCTASE) pir  S73840 ribonucleotide reductase 2 - Mycoplasma pneumoniae (strain ATCC 29342) gb AAB96162.1  (AE000050) ribonucleoside-diphosphate reductase beta chain [Mycoplasma pneumoniae] Length = 339	Identities = 105/309 (33%), Positives = 173/309 (55%), Gaps = 21/309 (6%) pir  T00087 rhamnosyltransferase - Streptococcus mutans dbj  BAA32090.1  (AB010970) rhamnosyltransferase [Streptococcus mutans]			Identities = 461/1194 (38%), Positives = 666/1194 (55%), Gaps = 74/1194 (6%) gb[AAB17762.1  (U56908) SCPB [Streptococcus agalactiae] Length = 1150	Identities = 65/143 (45%), Positives = 93/143 (64%), Gaps = 5/143 (3%) pir  H83632 conserved hypothetical protein PA0115 [imported] - Pseudomonas aeruginosa (strain PA01) gb AAG03505.1 AE004449_14 (AE004449) conserved hypothetical protein [Pseudomonas aeruginosa] Length = 150	Identities = 48/180 (26%), Positives = 84/180 (46%), Gaps = 4/180 (2%) pir[ F82497 uridine phosphorylase VCA0134 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF96047.1  (AE004354) uridine phosphorylase [Vibrio cholerae] Length = 243
92	87	56	No Hits found	No Hits found	53	09	32
Contig116 (18069-18470 m)	Contig116 (18483-19493 m)	Contig116 (20046-20954 m)	Contig116 (21204-21404 p)	Contig116 (24213-24722 p)	Contig116 (21164-24883 m)	Contig116 (25053-25481 m)	Contig116 (25515-26279 m)
SA-541.1	SA-542.1	SA-544.1	SA-545.2	SA-546.1	SA-547.1	SA-548.1	SA-549.1
SeqID 1913	SeqID 1914	SeqID 1915	SeqID 1916	SeqID 1917	SeqID 1918	SeqID 1919	SeqID 1920

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Identities = 64/104 (61%), Positives = 82/104 (78%) spiP55339IECSA_BACSU ABC-TYPE TRANSPORTER ATP-BINDING PROTEIN ECSA pir  F69619 ABC transporter (ATP-binding protein) ecsA - Bacillus subtilis emb CAA61074.1  (X87807) putative ATP-binding protein of ABC-type [Bacillus subtilis] emb CAA7409.1  (Y14077) Hypothetical protein [Bacillus subtilis] emb CAB12844.1  (Z99109) ABC transporter (ATP-binding protein) [Bacillus subtilis]	Identities = 216/448 (48%), Positives = 297/448 (66%), Gaps = 4/448 (0%) pir[ F69806 RNA methyltransferase homolog yfjO - Bacillus subtilis emb CAB12631.1  (299108) similar to RNA methyltransferase [Bacillus subtilis] dbj BAA24300.1  (D78509) YfjO [Bacillus subtilis]	Identities = 73/263 (27%), Positives = 140/263 (52%), Gaps = 9/263 (3%) dbj BAB04643.1  (AP001510) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 266	Identities = 96/175 (54%), Positives = 122/175 (68%) dbj BAB04659.1  (AP001510) unknown conserved protein in B. subtilis [Bacillus halodurans] Length = 175	Identities = 142/331 (42%), Positives = 204/331 (60%), Gaps = 2/331 (0%) gb AAF61315.1  (U96166) unknown [Streptococcus cristatus]	Identities = 66/194 (34%), Positives = 98/194 (50%), Gaps = 9/194 (4%) dbj BAA94320.1  (AB033763) hypothetical protein [Staphylococcus aureus]
. <b>77</b>	49	90	89	09	19
	Contig116 (26546-27901 m)	Contig116 (28099-28776 p)	Contig116 (28860-29393 p)	Contig121 (35263-36249 p)	Contig121 (33762-35270 p)
SA-55.1	SA-550.1	SA-551.1	SA-552.1	SA-554.1	SA-555.1
SeqiD 1921	SeqID 1922	SeqID 1923	SeqID 1924	SeqID 1925	SeqID 1926



SeqID 1927	SA-557.1	Contig121 (31361-33748 p)	63	Identities = 336/794 (42%), Positives = 507/794 (63%), Gaps = 29/794 (3%) splP28366 SEGA_BAGSU PREPROTEIN TRANSLOCASE SECA SUBUNIT pir  JQ0647 preprotein translocase secA - Bacillus subtilis dbj BAA01122.1  (D10279) secA protein [Bacillus subtilis] gb AAC44957.1  (U56901) involved in protein export [Bacillus subtilis] emb CAB15547.1  (299122) translocase binding subunit (ATPase) [Bacillus subtilis]
SeqID 1928	SA-558.1	Contig121 (30364-31374 p)	No Hits found	
SeqID 1929	SA-559.1	Contig121 (28826-30385 p)	No Hits found	
SeqID 1930	SA-56.1	Contig131 (56872-57270 p)	No Hits found	
SeqID 1931	SA-560.1	Contig121 (27275-28819 p)	No Hits found	
SeqID 1932	SA-561.1	Contig121 (26046-27275 p)	90	Identities = 105/422 (24%), Positives = 213/422 (49%), Gaps = 49/422 (11%) pir  C82917 preprotein translocase UU250 [imported] - Ureaplasma urealyticum gb AAF30659.1 AE002122_28 (AE002122) preprotein translocase [Ureaplasma urealyticum] Length = 471
SeqID 1933	SA-562.1	Contig121 (24726-25922 p)	10	Identities = 30/78 (38%), Positives = 42/78 (53%)   gb AAF61315.1  (U96166) unknown [Streptococcus cristatus]   Length = 442
SeqID 1934	SA-563.1	Contig121 (23790-24665 p)	43	dentities = 88/228 (38%), Positives = 136/228 (59%), Gaps = 10/228 (4%) gb AAC44016.1  (U40830) Epsl (Streptococcus thermophilus) prf  2209356K epsl gene [Streptococcus thermophilus]
SeqID 1935	SA-565.1	Contig121 (22613-23797 p)	35	Identities = 91/256 (35%), Positives = 146/256 (56%), Gaps = 8/256 (3%) gb AAF28363.1 AF224467_2 (AF224467) putative glycosyl transferase [Haemophilus ducreyi] Length = 269
SeqID 1936	SA-566.1	Contig121 (21382-22623 p)	37	Identities = 103/259 (39%), Positives = 156/259 (59%), Gaps = 3/259 (1%) gb AAF28363.1 AF224467_2 (AF224467) putative glycosyl transferase [Haemophilus ducreyi] Length = 269

SA=567:1 Contig121 (20180-21385 p) 38	Identities = 68/286 (23%), Positives = 122/286 (41%), Gaps = 18/286 (6%) splP37749JYEFG_ECOLI HYPOTHETICAL 37.8 KD PROTEIN IN GND-RFC INTERGENIC REGION (GALF TRANSFERASE) pir[ 169645 probable nucleotide sugar synthetiase   Escherichia coli   gb AAB8405.1   (U09876) putative Galf transferase   Escherichia coli   db BAA15876.1   (U09876) putative Galf transferase   Escherichia coli   db BAA15878.1   (D90841) ORF_ID:0351.10-similar to   SwissProt Accession Number P37749   Escherichia coli   gb AAC75095.1   (D90842) ORF_ID:0351.10; similar to   SwissProt Accession Number P37749   Escherichia coli   gb AAC75095.1   (AE000294) putative Galf transferase   Escherichia coli   K12   Length = 330	ldentities = 180/657 (27%), Positives = 283/657 (42%), Gaps = 31/657 (4%) emb CAB65343.1  (AJ007010) liver stage antigen-3 [Plasmodium falciparum] Length = 1786	7.1 Contig131 (56480-56863 p) No Hits found 0.1 Contig121 (15272-15448 m) No Hits found	<del>,</del>	Identities = 201/493 (40%), Positives = 302/493 (60%), Gaps =	Contig121 (13022-14518 m) 59
	SA-568.1	SA-569.1	SA-57.1 SA-570.1	SA-571.1	SA-572.1	_
SeqID 1937-	SeqID 1938	SeqID 1939	SeqID 1940 SeqID 1941	SeqID 1942	SeqID 1943	_



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Identities = 579/667 (86%), Positives = 635/667 (94%), Gaps = 1/667 (0%)·spl@54986 UVRB_STRPN EXCINUCLEASE ABC SUBUNIT B pir  A42385 excinuclease ABC chain B - Streptococcus pneumoniae plasmid pSB470 gb AAA27020.1  (M80215)-uvs402-protein [Streptococcus pneumoniae] Length = 668	Identities = 58/190 (30%), Positives = 96/190 (50%), Gaps = 11/190 (5%) pir  T34651 probable transmembrane protein - Streptomyces coelicolor emb CAA22372.1  (AL034446) putative transmembrane protein [Streptomyces coelicolor A3(2)] Length = 307	Identities = 109/195 (55%), Positives = 156/195 (79%), Gaps = 4/195 (2%) gb[AAF16724.1 AF141644_1 (AF141644) putative integral membrane protein [Lactococcus lactis]  191	Identities = 147/240 (61%), Positives = 192/240 (79%) ref[NP_069514.1] glutamine ABC transporter, ATP-binding protein (glnQ) [Archaeoglobus fulgidus] pir  H69334 glutamine transport protein glnQ - Archaeoglobus fulgidus gb AAB90561.1  (AE001058) glutamine ABC transporter, ATP-binding protein (glnQ) [Archaeoglobus fulgidus] Length = 242			Identities = 297/435 (68%), Positives = 345/435 (79%), Gaps = 7/435 (1%) spIP20964 OBG_BACSU SPOOB-ASSOCIATED GTP-BINDING PROTEIN pir  B32804 GTP-binding protein obg -Bacillus subtilis gb AAA22505.1  (M24537) GTP-binding protein [Bacillus subtilis] emb CAB14752.1  (Z99118) GTPase activity [Bacillus subtilis]
46	29	21	77	No Hits found	No Hits found	. 22
Contig121 (10934-12925 p)	Contig121 (10034-10873 p)	Contig121 (7587-9770 m)	Contig121 (6847-7587 m)	Contig121 (6348-6707 m)	Contig121 (6098-6256 p)	Contig121 (4759-6072 p)
SA-573.1	SA-574.1	SA-575.1	SA-576.1	SA-577.1	SA-578.1	SA-579.1
SeqID 1944	SeqID 1945	SeqID 1946	SeqID 1947	SeqID 1948	SeqID 1949	SeqID 1950

SeqID 1951	SA-58.1	Contig131 (55941-56345 p)	52	Identities = 25/79 (31%), Positives = 31/79 (38%) sp P04929 HRPX_PLALO HISTIDINE-RIGH-GL-YGOPROTEIN PRECURSOR pir  KGZQHL histidine-rich glycoprotein precursor Plasmodium lophurae emb CAA25698.1  (X01469) histidine-rich
				[Plasmodium sp.] Length = 351
SeqID 1952	SA-582.1	Contig121 (3151-4392 m)	73	Identities = 247/413 (59%), Positives = 314/413 (75%) spiQ9X4A7 PEPS_STRTR_AMINOPEPTIDASE PEPS gb AAD28348.1 AF102860_2 (AF102860) aminopeptidase PepS [Streptococcus thermophilus] Length = 413
SeqID 1953	SA-583.1	Contig121 (2613-2852 m)	No Hits found	
SeqID 1954	SA-584.1		34	Identities = 41/152 (26%), Positives = 75/152 (48%), Gaps = 4/152 (2%) emb CAB88235.1  (AL353012) hypothetical serinerich repeat protein [Schizosaccharomyces pombe] Length = 451
SeqID 1955	SA-585.2	Contig121 (387-2429 p)	38	Identities = 173/492 (35%), Positives = 270/492 (54%), Gaps = 38/492 (7%) gb AAK04264.1 AE006254_5 (AE006254) amidase [Lactococcus lactis subsp. lactis]
SeqID 1956	SA-586.2	Contig121 (3-302 p)	25	Identities = 39/89 (43%), Positives = 57/89 (63%), Gaps = 4/89 (4%) dbj BAB06992.1  (AP001518) 16S pseudouridylate synthase [Bacillus halodurans]
SeqID 1957	SA-589.2	Contig122 (32148-33029 m)	98	Identities = 230/291 (79%), Positives = 257/291 (88%) splQ07211[SCRK_STRMU FRUCTOKINASE dbj BAA02467.1  (D13175) fructokinase [Streptococcus mutans] Length = 293
SeqID 1958	SA-59.1	Contig131 (55843-55980 m)	No Hits found	
SeqID 1959	SA-590.1	_	82	Identities = 232/312 (74%), Positives = 262/312 (83%) splQ59935 MANA_STRMU MANNOSE-6-PHOSPHATE ISOMERASE (PHOSPHOMANNOSE ISOMERASE) (PMI) (PHOSPHOHEXOMUTASE) dbj BAA04021.1  (D16594) Mannosephosphate Isomerase (Streptococcus mutans) Length = 316



SeqID 1960	SA-5 <u>91.1</u>	Contig122 (28446-30974 m)	92	Identities = 503/843 (59%), Positives = 643/843 (75%), Gaps =
SeqID 1961	 SA-592.1	Contig122 (27313-28320 m)	. 54	Identities = 122/348 (35%), Positives = 188/348 (53%), Gaps = 32/348 (9%) pir] A81791 2-dehydro-3-deoxyphosphoheptonate aldolase (EC 4.1.2.15) NMA2180 [imported] - Neisseria meningitidis (group A strain Z2491) emb CAB85392.1  (AL162758) phospho-2-dehydr-3-deoxyheptonate aldolase [Neisseria meningitidis] Length = 351
SeqID 1962	SA-593.1	Contig122 (26929-27288 m)	73	Identities = 61/120 (50%), Positives = 91/120 (75%), Gaps = 1/120 (0%) gb[AAG22706.1] (AF276617) acyl carrier protein synthase; AcpS [Streptococcus pneumoniae] Length = 120
SeqID 1963	SA-594.1	Contig122 (25832-26932 m)	72	Identities = 227/366 (62%), Positives = 270/366 (73%)   gb AAD51027.1 AF171873_1 (AF171873) alanine racemase   Streptococcus pneumoniae
SeqID 1964	SA-597.1	Ccntig122 (24201-25739 m)	99	Identities = 248/530 (46%), Positives = 341/530 (63%), Gaps = 18/530 (3%) gb AAB52379.1  (U31811) immunogenic secreted protein precursor [Streptococcus pyogenes] Length = 534
SeqID 1965	SA-598.1	Contig122 (22106-24121 m)	84	Identities = 483/671 (71%), Positives = 568/671 (83%) splQ54900 RECG_STRPN ATP-DEPENDENT DNA HELICASE RECG pirl S71016 helicase recG homolog - Streptococcus pneumoniae emb CAA90280.1  (Z49988) MmsA [Streptococcus pneumoniae] prf  2209420A mmsA gene [Streptococcus pneumoniae]
SeqID 1966	SA-599.1	Contig122 (20901-21821 m)	72	Identities = 173/300 (57%), Positives = 224/300 (74%) dbj BAB07646.1  (AP001520) aryl-alcohol dehydrogenase [Bacillus halodurans] Length = 305

Identities = 31/81 (38%), Positives = 43/81 (52%), Gaps = 18/81 (22%) sp P43813 DNLJ_HAEIN DNA LIGASE (POLYDEOXYRIBONUCLEOTIDE SYNTHASE [NAD+]) pirl D64182 DNA ligase (NAD+) (EC 6.5.1.2) - Haemophilus influenzae (strain Rd KW20) gb AAC22753.1  (U32789) DNA ligase (lig) [Haemophilus influenzae Rd] Length = 679	Identities = 42/95 (44%), Positives = 56/95 (58%), Gaps = 4/95 (4%) ref[XP_001738.1] megakaryocyte stimulating factor [Homo sapiens] Length = 1385	Identities = 114/279 (40%), Positives = 171/279 (60%), Gaps = 3/279 (1%) splP28244 YDIB_ECOLI HYPOTHETICAL 31.2 KDA PROTEIN IN LPP-AROD INTERGENIC REGION pir  D64927 probable shikimate 5-dehydrogenase (EC 1.11.25) ydiB - Escherichia coli dbj BAA15449.1  (D90811) Shikimate 5-dehydrogenase (EC 1.1.1.25). [Escherichia coli] gb AAC74762.1  (AE000264) putative oxidoreductase [Escherichia coli K12]	Identities = 158/319 (49%), Positives = 214/319 (66%), Gaps = 4/319 (1%) dbj BAB05343.1  (AP001512) L-asparaginase [Bacilius halodurans] Length = 322	Identities = 89/281 (31%), Positives = 141/281 (49%), Gaps = 31/281 (11%) pir  C69862 conserved hypothetical protein ykrA - Bacillus subtilis emb  CAB13328.1  (Z99111) similar to hypothetical proteins [Bacillus subtilis] gb  AAC24929.1  (AF012285) unknown [Bacillus subtilis] Length = 257	Identities = 62/141 (43%), Positives = 93/141 (64%) dbj BAB06903.1  (AP001518) BH3184~unknown conserved protein [Bacillus halodurans] Length = 147
42	31	99	64	30	55
Contig137 (39066-39299 p)	Contig:131 (55778-56308 p)	Contig122 (19925-20803 m)	Contig122 (18928-19890 p)	Contig122 (17477-18859 m)	Contig122 (16969-17421 p)
SA-6.1	SA-60.1	SA-600.1	SA-601.1	SA-602.1	SA-603.2
SeqID 1967	SeqID 1968	SeqID 1969	SeqID 1970	SeqID 1971	SeqID 1972



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Identities = 250/404 (61%), Positives = 328/404 (80%), Gaps = 1/404 (0%) splP71348 YFBQ_HAEIN PROBABLE	AMINOTRANSFERASE HI0286 gb AAC21948.1  (U32714) aminotransferase [Haemophilus influenzae Rd] Length = 404	Identities = 129/257 (50%), Positives = 181/257 (70%), Gaps = 3/257 (1%) dbj BAB06181.1  (AP001515) transcriptional pleiotropic repressor [Bacillus halodurans] Length = 259	Identities = 101/183 (55%), Positives = 133/183 (72%) pir  C70008 pyrazinamidase/nicotinamidase homolog yueJ - Bacillus subtilis emb CAB15164.1  (299120) similar to pyrazinamidase/nicotinamidase [Bacillus subtilis] Length = 183	Identities = 151/321 (47%), Positives = 196/321 (61%), Gaps = 36/321 (11%) pirJB75610 probable 3-hydroxyacyl-CoA dehydrogenase - Deinococcus radiodurans (strain R1) gbJAAF12219.1JAE001862_45 (AE001862) 3-hydroxyacyl-CoA dehydrogenase, putative [Deinococcus radiodurans] Length = 347	Identities = 88/210 (41%), Positives = 141/210 (66%), Gaps = 3/210 (1%) pir[]A69969 conserved hypothetical protein yqzB - Bacillus subtilis emb[CAB14454.1] (Z99116) similar to hypothetical proteins [Bacillus subtilis] emb[CAB14467.1] (Z99117) similar to hypothetical proteins [Bacillus subtilis] Length = 212	Identities = 126/264 (47%), Positives = 186/264 (69%), Gaps = 1/264 (0%) dbj BAB05092.1  (AP001511) unknown conserved protein [Bacillus halodurans] Length = 270	Identities = 495/870 (56%), Positives = 648/870 (73%), Gaps = 4/870 (0%) splP22983jPODK_CLOSY PYRUVATE,PHOSPHATE DIKINASE (PYRUVATE,ORTHOPHOSPHATE DIKINASE) Length = 874
		62	, 69	51	64	99	73
	Contig122 (15490-16701 m)	Contig122 (14579-15364 m)	Contig122 (13964-14512 m)	Contig122 (12952-13917 p)	Contig122 (12134-12646 m)	Contig122 (11293-12123 m)	Contig122 (8635-11280 m)
	SA-604.2	SA-605.1	SA-606.1	SA-607.1	SA-608.1	SA-609.1	SA-611.1
	SeqID 1973	SeqID 1974	SeqID 1975	SeqID 1976	SeqID 1977	SeqID 1978	SeqID 1979

Identities = 42/96 (43%), Positives = 64/96 (65%)  dbjjBAB04384.1[(AP001509) glutamyl-tRNA (Gln) amidotransferase subunit C [Bacillus halodurans] Length = 96	Identities = 285/485 (58%), Positives = 365/485 (74%), Gaps = 2/485 (0%) splO06491 GATA_BACSU GLUTAMYL-TRNA(GLN) AMIDOTRANSFERASE SUBUNIT A (GLU-ADT SUBUNIT A) pir  B69795 glutamyl-tRNA(Gln) amidotransferase (EC 2.6) chain A [validated] - Bacillus subtilis emb CAB12488.1  (299107) alternate gene name: yedB~similar to amidase [Bacillus subtilis]	Identities = 309/476 (64%), Positives = 361/476 (74%), Gaps = 1/476 (0%) splQ9Z9X0jGATB_BACHD GLUTAMYL-TRNA(GLN) AMIDOTRANSFERASE SUBUNIT B (GLU-ADT SUBUNIT B) pir  T44293 hypothetical protein yerN [imported] - Bacillus halodurans dbj BAA75312.1  (AB011836) similar to B.subtilis yerN gene(87 identity) [Bacillus halodurans] dbj BAB04386.1  (AP001509) glutamyl-tRNA (Gln) amidotransferase subunit B [Bacillus halodurans] Length = 476	Identities = 102/292 (34%), Positives = 169/292 (56%), Gaps = 3/292 (1%) pir  T30575 hypothetical protein - Bacillus megaterium (fragment) emb CAA04271.1  (AJ000733) hypothetical protein [Bacillus megaterium]	Identities = 52/153 (33%), Positives = 88/153 (56%) spjP54452 YQEG_BACSU HYPOTHETICAL 20.1 KD PROTEIN IN NUCB-AROD INTERGENIC REGION pirj C69951 conserved hypothetical protein yqeG - Bacillus subtilis dbj BAA12443.1  (D84432) YqeG [Bacillus subtilis] emb CAB14510.1  (299117) similar to hypothetical proteins [Bacillus subtilis] Length = 172
. 19	£7	74	54	42
Contig122-(8195-8497 m)	Contig122 (6729-8195 m)	Contig122 (5287-6729 m)	Contig122 (4243-5157 m)	Contig122 (3599-4126 m)
SA-612.1	SA-613.1	SA-614.1	SA-615.1	SA-616.1
SeqID 1980	SeqID 1981	SeqID 1982	SeqID-1983	SeqID 1984



351 351	97	11	(+) (+)
%), Gap: AL 41.0 pir  D69! subtilis subtilis	3%), Gaps = 2/ FICAL 10.8 KD IIC REGION blog yqel - Bac acillus subtilis] drodipicolinate Length = 96	(47%), Gaps integrase Length = 362	187 (71%) COTINATE- SAMIDO-NAD- NAD(+) INATE RASE) (NAMN- rved hypothet (D84432) Yq 7) similar to
373 (74° THETIC EGION Bacıllus 9 Sillus sut	68%), G ETICAL ENIC RE nolog yc Bacillus iydrodip Lengtl	63 (47% 53) integ Leng	es = 134/187 (7 ABLE NICOTINA (ASE (DEAMIDC EAMIDO-NAD(+ (NICOTINATE (NICOTINATE (ANSFERASE) (51 CONSERVED hy (12447.11 (D844 11 (299117) simi
s = 280/373 (7 U HYPOTHET ENIC REGION yqeH - Bacıllus qeH [Bacillus s ar to hypothetic	67/97 (VYPOTHI VTERGE tase hor 2) Yqel   2) Yqel   ar to dilis]	= 126/2   (U404)   [12]	tives = 1 BABLE ERASE (DEAMII (NIC TRANSE 9951 cor AA1244
Positive BACSI NTERG protein 1432) Yc	sitives = 67 ACSU HYP DMER INTE te reductas (D84432) \(17) similar subtilis]	ositives 48867.1 phage	%), Posi SU PRC RANSFI (LASE) (SE) ENYLYL pir  F6( is abj B,
tities = 219/373 (58%). Positives = 280/373 (74%). Ga (2%) splP54453 YQEH_BACSU HYPOTHETICAL 41 TEIN IN NUCB-AROD-INTERGENIC REGION pir  D6 conserved hypothetical protein yqeH - Bacıllus subtilis db  BAA12444 1  (D84432) YqeH [Bacillus subtilis] ib  CAB14509 1  (299117) similar to hypothetical protei [Bacıllus subtilis]	itiles = 49/97 (50%), Positives = 67/97 (68%), Gaps = %) splP54454fYQEI_BACSU HYPOTHETICAL 10.8 K PROTEIN IN AROD-COMER INTERGENIC REGION 59951 dihydrodipicolinate reductase homolog yqel - Babtilis dbj BAA12446.1  (D84432) Yqel [Bacillus subtilinh]CAB14507.1  (Z99117) similar to dihydrodipicolinal reductase [Bacillus subtilis]	ities = 77/263 (29%), Positives = 126/263 (47%), Ga 34/263 (12%) gb AAC48867.1  (U40453) integrase eptococcus pyogenes phage T12]	Identities = 85/187 (45%), Positives = 134/187 (71%) sp P54455 NADD_BACSU PROBABLE NICOTINATE-CLEOTIDE ADENYLYLTRANSFERASE (DEAMIDO-NAIPROPHOSPHORYLASE) (DEAMIDO-NAD(+) DIPHOSPHORYLASE) (NICOTINATE ONOUCLEOTIDE ADENYLYLTRANSFERASE) (NAMINYLYLTRANSFERASE) pir  F69951 conserved hypotherin yqeJ - Bacillus subtilis dbj BAA12447.1  (D84432) Y [Bacillus subtilis] emb CAB14506.1  (299117) similar to hypothetical proteins [Bacillus subtilis] Length = 18
219/373 ppP5445 v NUCB ved hypi ved hypi AA1244 14509 1	ities = 49/97 (50%) %) splP54454 YQE  PROTEIN IN AROC 9951 dihydrodipico btilis dbj BAA1244  nb CAB14507.1  (Zindled)	77/263 33 (12%)	es = 85/ 455 NAC DE ADEI ROPHOS PHOSPI ICLEOT RANSF 1 - Bacilli s subtilis
Identities = 219/373 (58%). Positives = 280/373 (74%). Gaps = 8/373 (2%) splP54453 YOEH_BACSU HYPOTHETICAL 41.0 KD PROTEIN IN NUCB-AROD-INTERGENIC REGION pir  D69951 conserved hypothetical protein yqeH - Bacıllus subtilis db  BAA12444 11 (D84432) YqeH [Bacillus subtilis] emb  CAB14509 11 (299117) similar to hypothetical proteins [Bacillus subtilis]	Identities = 49/97 (50%), Positives = 67/97 (68%), Gaps = 2/97 (2%) splP54454 YQEI_BACSU HYPOTHETICAL 10.8 KD PROTEIN IN AROD-COMER INTERGENIC REGION pir  E69951 dihydrodipicolinate reductase homolog yqel - Bacillus subtilis dbj  BAA12446.1  (D84432) Yqel [Bacillus subtilis] emb  CAB14507.1  (299117) similar to dihydrodipicolinate reductase [Bacillus subtilis]	Identities = 77/263 (29%), Positives = 126/263 (47%), Gaps = 34/263 (12%) gb[AAC48867.1  (U40453) integrase [Streptococcus pyogenes phage T12] Length = 362	Identities = 85/187 (45%), Positives = 134/187 (71%) splP54455 NADD_BACSU PROBABLE NICOTINATE-NUCLEOTIDE ADENYLYLTRANSFERASE (DEAMIDO-NAD(+) PYROPHOSPHORYLASE) (DEAMIDO-NAD(+) DIPHOSPHORYLASE) (NICOTINATE MONONUCLEOTIDE ADENYLYLTRANSFERASE) (NAMN ADENYLYLTRANSFERASE) pirl F69951 conserved hypothetical protein yqeJ - Bacillus subtilis dbj BAA12447.1  (D84432) YqeJ  Bacillus subtilis] emb CAB14506.1  (299117) similar to hypothetical proteins [Bacillus subtilis] Length = 189
1de   8/37   PR(		<u>5</u>	N N O O O
73	26	44	9
,			
(m 6	(E 8	(m 20	1 m)
 (2481-3599 m)	(2071-2388 m)	55074-55907 m)	(1309-1941 m)
Contig 122 (2	Contig122 (2		Contig122 (1:
Cont	Cont	Contig131	Conti
SA-617.1	SA-618.1	SA-62.1	SA-620.1
		SA-	
SeqID 1985	SeqID 1986	SeqID 1987	SeqID.1988
Seq	Seq	Seq	Seq

			·	Identities = 79/180 (43%), Positives = 116/180 (63%) -sp P54456 YQEK_BACSU HYPOTHETICAL 21.3 KD PRGTEIN- IN AROD-COMER INTERGENIC REGION pir  G69951
SeqID 1989	SA-621.1	Contig122 (725-1312 m)	ග	conserved hypothetical protein yqeK - Bacillus subtilis ———dbj BAA12448.1  (D84432) YqeK [Bacillus subtilis] emb CAB14505.1  (Z99117) similar to hypothetical proteins [Bacillus subtilis] Length = 186
SeqID 1990	SA-622.1	Contig122 (140-682 m)	42	Identities = 46/175 (26%), Positives = 81/175 (46%), Gaps = 12/175 (6%) gb AAG19496.1  (AE005041) Vng1100c [Halobacterium sp. NRC-1]
SeqID 1991	SA-623.2	Contig129 (30428-31114 m)	52	Identities = 80/214 (37%), Positives = 122/214 (56%), Gaps = 3/214 (1%) gb AAC95438.1  (AF068901) unknown [Streptococcus pneumoniae] Length = 234
SeqID 1992	SA-624.1	Contig129 (31243-32463 m)	74	Identities = 220/398 (55%), Positives = 306/398 (76%), Gaps = 3/398 (0%) gb AAF36228.1 AF168363_4 (AF168363) oxalate:formate antiporter [Lactococcus lactis] Length = 421
SeqID 1993	SA-625.1	Contig129 (32650-34017 m)	76	Identities = 313/453 (69%), Positives = 375/453 (82%) gb AAC95436.1  (AF068901) D-Ala-D-Ala adding enzyme [Streptococcus pneumoniae] Length = 457
SeqID 1994	SA-626.1	Contig129 (34164-35210 m)	8	Identities = 243/346 (70%), Positives = 289/346 (83%) splO54631 DDL_STRPN D-ALANINED-ALANINE LIGASE (D-ALANYLALANINE SYNTHETASE) (D-ALA-D-ALA LIGASE) gb AAC95435.1  (AF068901) D-Ala-D-Ala ligase [Streptococcus pneumoniae] Length = 347
SeqID 1995	SA-627.1	Contig129 (35351-35947 m)	. 76	Identities = 181/198 (91%), Positives = 189/198 (95%) sp P96053 RECR_STRTR RECOMBINATION PROTEIN RECR gb AAC44615.1  (U58210) RecM [Streptococcus thermophilus] Length = 198
SeqID 1996	SA-628.1	Contig129 (35962-38004 m)		Identities = 346/698 (49%), Positives = 482/698 (68%), Gaps = 17/698 (2%) gbjAAC44614.1 (U58210) penicillin-binding protein 2b [Streptococcus thermophilus]



SeqID 1997	SA-629.1	Contig129 (38136-38828 m)	95	Identities = 219/230 (95%), Positives = 226/230 (98%) emb CAB51328:t -(AJ131985) phosphoglyceromutase [Streptococcus pneumoniae] Length = 230
SeqID 1998	SA-63.1	Contig131 (54046-54684 m)	91	Identities = 30/100 (30%), Positives = 43/100 (43%), Gaps = 21/100 (21%) gb AAG10259.1 AF264920_1 (AF264920) DS06238.4-like protein [Drosophila yakuba] Length = 213
SeqID 1999	SA-630.1	Contig129 (39005-39763 m)	2.2	Identities = 164/252 (65%), Positives = 202/252 (80%) splP50918 TPIS_LACLA TRIOSEPHOSPHATE ISOMERASE (TIM) gb AAC43268.1  (U07640) triosephosphate isomerase [Lactococcus lactis] Length = 252
SeqID 2000	SA-632.1	Contig129 (40284-40760 p)	45	Identities = 48/145 (33%), Positives = 73/145 (50%), Gaps = 6/145 (4%) pir[ T39903 serine-rich protein - fission yeast (Schizosaccharomyces pombe) emb[CAA22127.1  (AL033534) hypothetical serine-rich secreted protein [Schizosaccharomyces pombe] Length = 534
SeqID 2001	SA-633.1	Contig129 (39944-41140 m)	94	Identities = 384/398 (96%), Positives = 396/398 (99%) splP33170jEFTU_STROR ELONGATION FACTOR TU (EF-TU) pirjF60663 translation elongation factor EF-Tu - Streptococcus oralis Length = 398
SeqID 2002	SA-635.1	Contig129 (41492-42709 m)	53	Identities = 149/423 (35%), Positives = 229/423 (53%), Gaps = 41/423 (9%) sp[Q47866 FTSW_ENTHR PROBABLE CELL DIVISION PROTEIN FTSW gb AAB39929.1  (U58049) putative cell division protein ftsW [Enterococcus hirae] Length = 397
SeqID 2003	SA-636.1	Contig129 (42866-45661 m)	59	Identities = 362/945 (38%), Positives = 559/945 (58%), Gaps = 65/945 (6%) splO32483 CAPP_RHOPA PHOSPHOENOLPYRUVATE CARBOXYLASE (PEPCASE) (PEPC) dbj BAA21835.1  (D89668) phosphoenolpyruvate carboxylase [Rhodopseudomonas palustris] Length = 936
SeqID 2004	SA-637.1	Contig129 (45870-47669 p)	80	Identities = 443/519 (85%), Positives = 489/519 (93%) dbj BAB16034.1  (AB030810) Bacillus licheniformis Pz-peptidase homologue [Streptococcus pyogenes] Length = 519



puno	puno	Identities = 102/221 (46%), Positives = 145/221 (65%)   emb CAC07978.1  (AJ278983) CopR protein [Ralstonia   metallidurans] Length = 228	Identities = 64/210 (30%), Positives = 95/210 (44%), Gaps = 15/210 (7%) emb CAA72266.1  (Y11477) endolysin [Bacteriophage Bastille] Length = 364	Identities = 38/156 (24%), Positives = 83/156 (52%), Gaps = 13/156 (8%) gb AAG20117.1  (AE005090) NADH dehydrogenase/oxidoreductase-like protein; NolA [Halobacterium sp. NRC-1] Length = 303	Identities = 61/164 (37%), Positives = 96/164 (58%), Gaps = 13/164 (7%) splP36922 EBSC_ENTFA EBSC PROTEIN pir  C49939 ebsC protein - Enterococcus faecalis gb AAC36853.1  (L23802) regulatory protein [Enterococcus faecalis] Length = 164	Identities = 65/191 (34%), Positives = 93/191 (48%), Gaps = 13/191 (6%) pir  G72260 phosphoglycerate mutase - Thermotoga maritima (strain MSB8) gb AAD36444.1 AE001791_6 (AE001791) phosphoglycerate mutase [Thermotoga maritima] Length = 201	Identities = 69/232 (29%), Positives = 108/232 (45%), Gaps = 9/232 (3%) pir[ E69814 conserved hypothetical protein yfnB - Bacillus subtilis db  BAA20111.1  (D86418) YfnB (Bacillus subtilis) emb  CAB12552.1  (Z99107) similar to hypothetical proteins [Bacillus subtilis] emb  CAB12562.1  (Z99108) similar to hypothetical proteins [Bacillus subtilis] Length = 235
No Hits found	No Hits found	61	32	98	69	42	
1 -	Contig129 (48101-48571 m)	Contig131 (53290-53961 m)	Contig129 (48875-49630 p)	Contig129 (49663-50280 m)	Contig129 (50575-51048 p)	Contig129 (51058-51711 p)	Contig129 (51747-52649 m)
SA-638.1	SA-639.1	SA-64.1	SA-640.1	SA-641.1	SA-642.1	SA-643.1	SA-644.1
SeqID 2005	SegID 2006	SeqID 2007	SeqID 2008	SeqID 2009	SeqID 2010	SeqID 2011	Seq1D 2012



				Identities = 318/491 (64%), Positives = 391/491 (78%), Gaps =
SeqID 2013	 SA-646.1	Contig 129 (52812-54314 p)	20	SYNTHETASE (LYSINE-TRNA LIGASE) (LYSRS) pirils66111 lysine-tRNA ligase (EC 6.1.1.6) lysS - Bacillus subtilis
			•	dbj BAA053:16-1  (D26185) lysyl-tRNA thynthetase [Bacillus subtilis] emblCAB11858.11 (Z99104) lysyl-tRNA synthetase
				[Bacillus subtilis] Length = 499
				Identities = 103/151 (68%), Positives = 120/151 (79%)
				SPIP 11996 RISB_BACSU 6,7-DIMETHYL-8-RIBITYLLUMAZINE   SYNTHASE (DMRL SYNTHASE) (LUMAZINE
				SYNTHASE) (RIBOFLAVIN SYNTHASE BETA CHAIN)
				pir  A26708 riboflavin synthase (EC 2.5.1.9) complex beta chain
				ribH [validated] - Bacillus subtilis pdb/1RW/1 Chain 1,
				SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF BACILLUS
				SUBTILIS pdb/1RVV/2 Chain 2, SynthaseRIBOFLAVIN
				SYNTHASE COMPLEX OF BACILLUS SUBTILIS pdb 1RVV 3
				Chain 3, SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF
SealD 2014	SA-647 1	Contin 129 (54389-54859 m)	9	BACILLUS SUBTILIS pdb 1RVV 4 Chain 4,
	· · · · · · · · · · · · · · · · · · ·	( 0000000000000000000000000000000000	3	SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF BACILLUS
				SUBTILIS pdb[1RVV A Chain A, SynthaseRIBOFLAVIN
				SYNTHASE COMPLEX OF BACILLUS SUBTILIS pdb/1RVVJB
				Chain B, SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF
				BACILLUS SUBTILIS pobj1RVVJC Chain C,
				SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF BACILLUS
				SUBTILIS pdb/1RVV/D Chain D, SynthaseRIBOFLAVIN
				SYNTHASE COMPLEX OF BACILLUS SUBTILIS publication
				Chain E, SynthaseRIBOFLAVIN SYNTHASE COMPLEX OF
				BACILLUS SUBTILIS pdb/1RVV/F Chain F,
				SynthaseRIBOFLAVIN SYNTHA

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SeqID 2015	SA-648.1	Contig129 (54874-56067 m)		Identities = 230/395 (58%), Positives = 307/395 (77%) splP50855 GCH2_ACTPL_RIBOFLAVIN_BIOSYNTHESIS PROTEIN RIBA [INCLUDES: GTP CYCLOHYDROLASE : 3,4-DIHYDROXY-2-BUTANONE 4-PHOSPHATE SYNTHASE (DHBP SYNTHASE)] :pir  †750548 GTP cyclohydrolase II (EC 3.5.4.25) / 3, 4-dihydroxy-2-butanone 4-phosphate synthase (EC 5.4.99) [validated] - Actinobacillus pleuropneumoniae gb AAA86524.1  (U27202) GTP cyclohydrase II/ 3,4-dihydroxy-2-butanone-4-phosphate synthase [Actinobacillus pleuropneumoniae] Length = 401
SeqID 2016	SA-649.1	Contig129 (56085-56735 m)	29	Identities = 98/216 (45%), Positives = 147/216 (67%), Gaps = 2/216 (0%) dbj BAB05274.1  (AP001512) riboflavin synthase alpha subunit [Bacillus halodurans] Length = 215
SeqID 2017	SA-65.1	Contig131 (51942-53309 m)	49.	Identities = 114/466 (24%), Positives = 227/466 (48%), Gaps = 42/466 (9%) dbj BAB04092.1  (AP001508) two-component sensor histidine kinase [Bacillus halodurans] Length = 459
SeqID 2018	SA-651.1	Contig129 (56716-57825 m)	99	Identities = 180/353 (50%), Positives = 257/353 (71%) spiP50853 RIBD_ACTPL RIBOFLAVIN BIOSYNTHESIS PROTEIN RIBD [INCLUDES: DIAMINOHYDROXYPHOSPHORIBOSYLAMINOPYRIMIDINE DEAMINASE (RIBOFLAVIN-SPECIFIC DEAMINASE); 5-AMINO-6-(5-PHOSPHORIBOSYLAMINO)URACIL REDUCTASE (HTP REDUCTASE)] Length = 376
SeqID 2019	SA-653.1	Contig129 (58686-59894 p)		Identities = 192/436 (44%), Positives = 274/436 (62%), Gaps = 14/436 (3%) pir  C82733 manganese transport protein XF1015 [imported] - Xylella fastidiosa (strain 9a5c) gb AAF83825.1 AE003939_4 (AE003939) manganese transport protein [Xylella fastidiosa] Length = 472



:				Identities = 63/215 (29%), Positives = 108/215 (49%), Gaps =
SA-	SA-654.2	Contig129 (59996-60805 m)	40	13/215 (6%) pirj H75355 hypothetical protein - Deinococcus radiodurans (strain R1) gb AAF11325.1 AE002018_7 (AE002018) hypothetical protein [Deinococcus radiodurans] Length = 250
S I	SA-655.2	Contig129 (60917-61129 m)	53	Identities = 28/64 (43%), Positives = 41/64 (63%) splO83371 Y352_TREPA HYPOTHETICAL PROTEIN TP0352 pir  F71333 hypothetical protein TP0352 - syphilis spirochete gb AAC65352.1  (AE001215) T. pallidum predicted coding region TP0352 [Treponema pallidum] Length = 85
Ś	SA-656.2	Contig129 (61256-62542 m)	62	Identities = 201/407 (49%), Positives = 277/407 (67%), Gaps = 2/407 (0%) pir  H69979 proteinase homolog yrrO - Bacillus subtilis emb CAB14676.1  (Z99117) similar to protease [Bacillus subtilis] Length = 422
S I	SA-657.1	Contig129 (62672-63598 m)	90	Identities = 92/304 (30%), Positives = 161/304 (52%), Gaps = 5/304 (1%) pir  G69979 proteinase homolog yrrN - Bacillus subtilis emb CAB14677.1  (299117) similar to protease [Bacillus subtilis] Length = 309
Ś	SA-658.2	Contig133 (43969-46152 m)	46	Identities = 294/409 (71%), Positives = 342/409 (82%), Gaps = 7/409 (1%) gb AAD00281.1  (U78600) putative ptsG protein [Streptococcus mutans]
S)	SA-659.1	Contig133 (43486-43767 p)	No Hits found	
()	SA-66.1	Contig131 (51118-51453 m)	84	Identities = 45/76 (59%), Positives = 54/76 (70%) pir  T44087 hypothetical protein [imported] - Staphylococcus aureus (fragment) Length = 151
(ŠI	SA-660.1	Contig133 (43100-43915 m)	No Hits found	
S I	SA-661.1	Contig133 (42071-42823 m)	1.2	Identities = 135/242 (55%), Positives = 183/242 (74%) emb CAC10170.1  (AJ278301) response regulator [Streptococcus pneumoniae]
S I	SA-662.1	Contig133 (41703-42002 m)	65	Identities = 44/88 (50%), Positives = 65/88 (73%) emb[CAC10175.1  (AJ278302) histidine kinase [Streptococcus pneumoniae) Length = 446

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Identities = 105/291 (36%), Positives = 161/291 (55%), Gaps = 18/291 (6%)-emb CAB88481.1  (AL353816) putative ABC transport system ATP-binding protein   Streptomyces coelicolor A3(2)  Length = 289		Identities = 78/262 (29%), Positives = 129/262 (48%), Gaps = 29/262 (11%) gb AAF71283.1 AF253562_7 (AF253562) racemase [Enterococcus faecalis] Length = 711		Identities = 49/164 (29%), Positives = 96/164 (57%), Gaps = 8/164 (4%) splP37081JPTRB_KLEPN PTS SYSTEM, SORBOSE-SPECIFIC IIB COMPONENT (EIIB-SOR) (SORBOSE-PERMEASE IIB COMPONENT) (PHOSPHOTRANSFERASE ENZYME II, B COMPONENT) (EIII-B-SOR) pir[IS50188 phosphotransferase system enzyme II component B, sorbose-specific Klebsiella pneumoniae emb[CAA46858.1] (X66059) EIII-B Sor PTS [Klebsiella pneumoniae] prf[I2022173D sorB gene [Klebsiella pneumoniae] Length = 164	Identities = 303/506 (59%), Positives = 389/506 (75%), Gaps = 6/506 (1%) splP44023 YFCC_HAEIN HYPOTHETICAL PROTEIN HI0594 pir  E64010 hypothetical protein HI0594 - Haemophilus influenzae (strain Rd KW20) gb AAC22251.1  (U32741) conserved hypothetical transmembrane protein [Haemophilus influenzae Rd] Length = 509	Identitles = 70/251 (27%), Positives = 132/251 (51%), Gaps = 6/251 (2%) gb AAC44680.1  (U65015) PTS permease for mannose subunit IIPMan [Vibrio furnissii] Length = 258	
55	No Hits found No Hits found	36	No Hits found No Hits found	55	74	48	
Cont <u>ig133 (40732-4</u> 1580 m)	Contig133 (39572-40726 m) Contig133 (39693-39929 p)	Contig133 (38309-39376 m)	Contig133 (37637-38014 m) Contig133 (37144-37584 p)	Contig133 (37128-37619 m)	Contig131 (50330-51832 p)	Contig133 (36303-37112 m)	
SA-663.1	SA-664.1 SA-665.2	. SA-666.1	SA-667.1 SA-668.1	SA-669.1	SA-67.1	SA-670.1	
SeqID 2030	SeqID 2031 SeqID 2032	SeqID 2033	SeqID 2034 SeqID 2035	SeqID 2036	SeqID 2037	SeqID 2038	



SeqID 2039	SA-671.1	Contig133 (35479-36306 m)		Identities = 94/280 (33%), Positives = 156/280 (55%), Gaps = 13/280 (4%) sp P08188 PTND_ECOLI PTS SYSTEM, MANNOSE-SPECIFIC IID COMPONENT (EIID-MAN) (MANNOSE-PERMEASE IID COMPONENT) (PHOSPHOTRANSFERASEENZYME II, D COMPONENT) (EII-M-MAN) pir  WQECMM phosphotransferase system enzyme II (EC 2.7.1.69), mannose-specific, factor IID - Escherichia coli gb AAA24445.1  (J02699) mannose permease subunit II-M-Man (Escherichia coli) dbj BAA15631.1  (D90826) PTS system, Mannose-specific IID component) (EIID-MAN) (Mannose-permease IID component) (EII-M-MAN) (Escherichia coli] gb AAC74889.1  (AE000276) PTS enzyme IID, mannose-specific [Escherichia coli K12]
SeqID 2040	SA-672.1	Contig133 (33692-35341 m)	24	Identities = 84/258 (32%), Positives = 138/258 (52%), Gaps = 23/258 (8%) dbj BAB05628.1  (AP001513) two-component sensor histidine kinase [Bacillus halodurans] Length = 597
SeqID 2041	SA-673.1	Contig133 (32915-33688 m)	99	Identities = 86/252 (34%), Positives = 147/252 (58%), Gaps = 17/252 (6%) emb CAB54580.1  (AJ006398) response regulator   Streptococcus pneumoniae  gb AAF31452.1 AF221126_1 (AF221126) putative response regulator [Streptococcus pneumoniae] Length = 245
SeqID 2042	SA-674.1	Contig133 (31866-32906 m)	53	Identities = 114/331 (34%), Positives = 184/331 (55%), Gaps = 3/331 (0%) gb[AAD18094.2] (U75349) periplasmic-iron-binding protein BitA [Brachyspira hyodysenteriae] Length = 336
SeqID 2043	SA-675.1	Contig133 (31147-31644 p)	46	Identities = 55/150 (36%), Positives = 84/150 (55%), Gaps = 2/150 (1%) pir  F75336 conserved hypothetical protein - Deinococcus radiodurans (strain R1) gb AAF11472.1 AE002031_1 (AE002031) conserved hypothetical protein [Deinococcus radiodurans] Length = 190

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SeqID 2044	SA-676.1	Contig133 (30110-31147 p)	02	Identities = 212/345 (61%), Positives = 268/345 (77%), Gaps = -1/345-(0%)-pir  C69830 glucanase homolog yhfE - Bacillussubtilis emb CAA74526.1  (Y14083) hypothetical protein [Bacillus subtilis] emb CAB12860.1  (Z99109) similar to glucanase  Bacillus subtilis] Length = 346
Seq1D 2045	SA-677.1	Contig133 (29644-30099 p)	80	Identities = 103/142 (72%), Positives = 123/142 (86%) emb[CAB70606.1  (Y18363) ribonucleotide reductase-like (Nrd-like) protein [Streptococcus dysgalactiae subsp. equisimilis] Length = 142
SeqID 2046	SA-678.1	Contig133 (27085-29487 p)	61	Identities = 398/694 (57%), Positives = 493/694 (70%), Gaps = 30/694 (4%) emb CAB70615.1  (AJ133440) cyclo-nucleotide phosphodiesterase, putative [Streptococcus dysgalactiae subsp. equisimilis] Length = 683
SeqID 2047	SA-68.1	Contig131 (49259-50185 p)	65	Identities = 162/313 (51%), Positives = 207/313 (65%), Gaps = 7/313 (2%) splQ46171 ARCC_CLOPE CARBAMATE KINASE emb CAA66367.1  (X97768) carbamate kinase [Clostridium perfringens]
SeqID 2048	SA-680.1	Contig133 (24609-26825 m)	63	Identities = 647/739 (87%), Positives = 696/739 (93%), Gaps = 1/739 (0%) splQ54089JRELA_STREQ PUTATIVE GTP PYROPHOSPHOKINASE (ATP:GTP 3 - PYROPHOSPHOTRANSFERASE) (PPGPP SYNTHETASE I) ((P)PPGPP SYNTHETASE) (STRINGENT RESPONSE-LIKE PROTEIN) pirJS39975 stringent response-like protein - Streptococcus equisimilis embJCAA51353.1 (X72832) stringent response-like protein [Streptococcus equisimilis] tength = 739
SeqID 2049	SA-681.1	Contig133 (24156-24599 m)	83	Identities = 104/145 (71%), Positives = 126/145 (86%) pir  S39974 hypothetical protein - Streptococcus equisimilis emb  CA451352.1  (X72832) ORF1 [Streptococcus equisimilis] prf  2009358D rel upstream ORF [Streptococcus equisimilis] Length = 147



SeqID 2050	SA-682.1	Contig133 (22978-23901 m)	70	Identities = 180/304 (59%), Positives = 227/304 (74%), Gaps = 4/304 (1%) pir[I746757 tipoprotein-Imb-[validated] - Streptococcus agalactiae gb AAD13796.1 (AF062533) Lmb [Streptococcus agalactiae] Length = 306
SeqID 2051	SA-686.1	Contig133 (21506-22951 m)	55	Identities = 213/463 (46%), Positives = 277/463 (59%), Gaps = 41/463 (8%) pir  T46758 hypothetical 92.4K protein - Streptococcus agalactiae gb AAD13797.1  (AF062533) unknown [Streptococcus agalactiae] Length = 822
SeqID 2052	SA-687.1	Contig133 (20370-21608 m)	24	Identities = 66/253 (26%), Positives = 101/253 (39%), Gaps = 64/253 (25%) pir  T46758 hypothetical 92.4K protein - Streptococcus agalactiae gb AAD13797.1  (AF062533) unknown [Streptococcus agalactiae] Length = 822
SeqID 2053	SA-688.2	Contig133 (19312-20088 p)	89	Identities = 139/255 (54%), Positives = 189/255 (73%), Gaps = 6/255 (2%) gbjAAA25174.1 (M35375) lactose repressor (lacR; alt.) [Lactococcus lactis] gbjAAA25176.1 (M60447) repressor protein [Lactococcus lactis] gbjAAA25186.1 (M60673) lacR [Lactococcus lactis]
SeqID 2054	SA-689.2	Contig139 (185493-186197 p)	59	Identities = 99/228 (43%), Positives = 149/228 (64%), Gaps = 3/228 (1%) pir  140084 gtcR protein - Bacillus brevis emb CAA55264.1  (X78502) gtcR [Brevibacillus brevis] Length = 242
SeqID 2055	SA-69.1	Contig131 (48249-49247 p)	87	Identities = 264/332 (79%), Positives = 292/332 (87%) emb[CAB75986.1  (AJ272085) ornithine carbamoyltransferase [Staphylococcus aureus] emb[CAB75987.1  (AJ272086) ornithine carbamoyltransferase Otc6850 [Staphylococcus aureus] Length = 333
SeqID 2056	SA-690.1	Contig139 (183328-185271 p)	2.2	Identities = 412/638 (64%), Positives = 506/638 (78%), Gaps = 7/638 (1%) dbj BAB06860.1  (AP001517) threonyl-tRNA synthetase 1 [Bacillus halodurans] Length = 645
SeqID 2057	SA-691.1	Contig139 (181537-182871 p)	52	Identities = 189/290 (65%), Positives = 234/290 (80%) emb CAA72250.1  (Y11463) ORF5 [Streptococcus pneumoniae] Length = 290

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SeqID 2058—	SA-692:1	Contig139 (180537-181535 p)	53	Identities = 120/350 (34%), Positives = 180/350 (51%), Gaps = 29/350 (8%) gb AAG19110.1  (AE005009) Vng0600c
SeqID 2059	SA-693.1	Contig139 (179026-180492 p)	92	Identities = 305/483 (63%), Positives = 378/483 (78%).   gb AAC35010.1  (AF055987) intracellular a-amylase   Streptococcus mutans]   Length = 486
SeqID 2060	SA-694.2	Contig139 (177890-178894 p)	94	Identities = 304/332 (91%), Positives = 320/332 (95%) dbj BAA88121.3  (AB028599) catabolite control protein A [Streptococcus bovis] Length = 334
SeqID 2061	SA-696.2	Contig139 (176595-177680 m)	81	Identities = 257/359 (71%), Positives = 304/359 (84%) gb AAC46293.1  (AF014460) PepQ [Streptococcus mutans] Length = 359
SeqID 2062	SA-697.1	Contig139 (174747-176537 p)	42	Identities = 148/586 (25%), Positives = 252/586 (42%), Gaps = 77/586 (13%) emb CAB66297.1  (AL136519) beta-N-acety glucosaminidase. [Streptomyces coelicolor A3(2)] Length = 615
SeqID 2063	SA-698.1	Contig139 (173973-174731 p)	No Hits found	
SeqID 2064	SA-699.1	Contig139 (172927-173766 p)	92	Identities = 197/271 (72%), Positives = 228/271 (83%) sp P44481 Y048_HAEIN PUTATIVE OXIDOREDUCTASE HI0048 pir  B64045 D-arabinitol 2-dehydrogenase homolog - Haemophilus influenzae (strain Rd KW20) gb AAC21726.1  (U32690) oxidoreductase [Haemophilus influenzae Rd] Length = 285
SeqID 2065	SA-7.1	Contig137 (38846-39100 m)	No Hits found	
SeqID 2066	SA-70.1	Contig131 (46851-48146 m)	44	Identities = 118/427 (27%), Positives = 199/427 (45%), Gaps = 25/427 (5%) dbjjBAB06435.1  (AP001516) two-component sensor histidine kinase [Bacillus halodurans] Length = 437
SeqID 2067	SA-700.1	Contig139 (171757-172803 p)	73	Identities = 203/343 (59%), Positives = 261/343 (75%) dbj BAB04425.1  (AP001509) D-mannonate dehydrolase [Bacillus halodurans] Length = 345



SeqID 2068	SA-701.1	Contig 139 (170339-171739 p)	62	Identities = 215/465 (46%), Positives = 295/465 (63%), Gaps = 7/465 (1%) db;IBAB04424 1  (AP001509) uronate isomerase [Bacillus halodurans] Length = 472
SeqiD 2069	SA-702.1	Contig139 (169705-170322 p)	51	Identities = 92/199 (46%). Positives = 124/199 (62%), Gaps = 6/199 (3%) prillF72422 KHG-KDPG bifunctional aldolase TM0066 [similarity] - Thermotoga maritima (strain MSB8) gb AAD35160 1 AE001693_6 (AE001693) 2-dehydro-3-deoxyphosphogluconate aldolase/4-hydroxy-2-oxoglutarate aldolase [Thermotoga maritima] Length = 205
SeqID 2070	SA-704.1	Contig139 (168917-169588 p)	50	Identities = 67/225 (29%), Positives = 119/225 (52%), Gaps = 17/225 (7%) splP42239JYCBG_BACSU HYPOTHETICAL TRANSCRIPTIONAL REGULATOR IN GLTP-CWLJ INTERGENIC REGION (ORF6) pirl B69753 transcription regulator GntR family homology cbG - Bacillus subtilis dbj BAA06471.1  (D30808) homologue of protein A which gene locates upstream pyruvate dehydrogenase gene cluster [Bacillus subtilis] emb CAB12044.1  (Z99105) similar to transcriptional regulator (GntR family) [Bacillus subtilis]

Identities = 103/200 (51%). Positives = 136/200 (67%), Gaps = 4/200 (2%) spl003158INUCE_STRPN DNA-ENTRY NUCLEASE (COMPETENCE-SPECIFIC NUCLEASE) pir[ S10641 endAprotein - Streptococcus pneumoniae emb CAA38134.1  (X54225) membrane nuclease (Streptococcus pneumoniae)	Identities = 151/256 (58%). Positives = 186/256 (71%), Gaps = 15/256 (5%) prr  S72385 hypothetical protein 9 - Enterococcus faecalis plasmid pAD1 emb CAA65668.1  (X96977) orf9 [Enterococcus faecalis] Length = 278	Identities = 139/236 (58%), Positives = 178/236 (74%) gb AAD45529.1 AF162655_1 (AF162655) choline transporter [Streptococcus pneumoniae] Length = 242	Identities = 36/142 (25%), Positives = 62/142 (43%), Gaps = 7/142 (4%) pirj D70463 hypothetical protein aq_1894 - Aquifex aeolicus gb AAC07706.1  (AE000762) putative protein [Aquifex aeolicus]	Identities = 89/267 (33%), Positives = 157/267 (58%), Gaps = 15/267 (5%) gb AAF27301.1 AF154674_2 (AF154674) ParA [Lactococcus lactis subsp. lactis] gb AAF27325.1  (AF178424) ParA [Lactococcus lactis]		Identities = 33/93 (35%), Positives = 53/93 (56%), Gaps = 2/93 (2%) ref[NP_038304.1  ORF4 [Streptococcus thermophilus bacteriophage 7201] gb[AAF26603.1 AF118440_1 (AF145054) ORF4 [Streptococcus thermophilus bacteriophage 7201] Length = 268			Identities = 106/222 (47%), Positives = 162/222 (72%), Gaps = 2/222 (0%) gb AG28336.1  (U88582) SatD [Streptococcus mutans] Length = 222
20	40	72	24	53	No Hits found	04	No Hits found	No Hits found	99
Contig139 (156980-157705 p)	Contig125 (39398-40735 m)	Contig131 (45077-45799 m)	Contig125 (38800-39384 m)	Contig125 (37843-38661 m)	Contig125 (37568-37846 m)	Contig125 (37172-37561 m)	Contig125 (36856-37167 m)	Contig125 (35394-36722 m)	Contig125 (34305-35003 m)
SA-717.2	SA-719.1	SA-72.1	SA-720.1	SA-721.1	SA-722.1	SA-723.1	SA-725.1	SA-726.1	SA-727.1
SeqID 2082	SeqID 2083	SeqID 2084	SeqID 2085	SeqID 2086	SeqID 2087	SeqID 2088	SeqID 2089	SeqID 2090	SeqID 2091



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Identities = 54/103 (52%), Positives = 70/103 (67%), Gaps = 2/103 (1%) gb AAG28337.1  (U88582) SatE [Streptococcus mutans] Length = 108	Identities = 346/521 (66%), Positives = 411/521 (78%), Gaps = 24/521 (4%) gb AAD17886.1  (AF100456), hyaluronate-associated protein precursor [Streptococcus equi]		Identities = 119/148 (80%), Positives = 136/148 (91%)   gb AAC17173.1  (AF065141) unknown [Streptococcus mutans]   Length = 356	Identities = 188/218 (86%), Positives = 205/218 (93%)   gb AAC17173.1  (AF065141) unknown [Streptococcus mutans]   Length = 356	172/320 (53%), Gaps SSU HYPOTHETICAL E IN IDH 3 REGION istidine kinase homolog AA03301.11 (D14399) s] emb[CAB16001.1] sor histidine kinase [Yxsngth = 325	Identities = 95/226 (42%), Positives = 136/226 (60%), Gaps = 10/226 (4%) pir  D70032 two-component response regulator [YvcQ] homolog yvcP - Bacillus subtilis emb CAB08062.1  (294043) hypothetical protein [Bacillus subtilis] emb CAB15477.1  (299121) similar to two-component response regulator [YvcQ] [Bacillus subtilis] Length = 237	Identities = 154/666 (23%), Positives = 300/666 (44%), Gaps = 40/666 (6%) gblAAF99695.1 AF267498_5 (AF267498) permease OrfY [Streptococcus mutans] Length = 640
26		No Hits found	. 83	68	95		45
Contig125 (33542-34177 m)	Contig125 (31914-33509 p)	Contig131 (44407-44805 p)	Contig125 (31012-31461 m)	Contig125 (30392-31078 m)	Contig125 (29462-30349 p)	Contig125 (28746-29414 p)	Contig125 (26819-28636 p)
SA-728.1	SA-729.1	SA-73.1	SA-730.1	SA-731.1	SA-732.1	SA-733.1	SA-734.1
SeqID 2092	SeqID 2093	SeqID 2094:	SeqID 2095	SeqID 2096	SeqID 2097	SeqiD 2098	SeqID 2099

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Identities = 118/242 (48%), Positives = 175/242 (71%), Gaps = 1/2627-26679 p) 68 1/242 (0%)-gb AAF99694.1 AF267498_4 (AF267498) ABC - transporter OrfX [Streptococcus mutans] Length = 246	Identities.= 59/146 (40%), Positives = 87/146 (59%), Gaps = 3/146 (2%) gb AAB08491.1  (U25181) nisin-resistance protein   (Lactococcus lactis)   Length = 318	Identities = 51/177 (28%), Positives = 104/177 (57%), Gaps = 5/177 (2%) gb AAB08491.1  (U25181) nisin-resistance protein   [Lactococcus lactis]	24286-24774 m) No Hits found	23406-24254 p) 62. 14790 (6%) emb CAB59827.1  (AJ012388) hypothetical protein [Lactococcus lactis] Length = 286	Identities = 326/506 (64%), Positives = 410/506 (80%), Gaps = 1/506 (0%) pir  T44634 choline transporter [imported] - Streptococcus pneumoniae gb AAD45530.1 AF162656_1 (AF162656) choline transporter [Streptococcus pneumoniae] (AF162656) choline transporter [Streptococcus pneumoniae]	22530-23273 p) No Hits found	21118-22452 p)  79  6mb CAB13486.1  (AJ000975) Gid protein Broilius subtilis] emb CAB13486.1  (A99112) glucose-inhibited division protein [Bacillus subtilis] emb CAB13486.1  (Bacillus subtilis] [Bacillus subtilis]	20306-21004 p) 56 1/229 (0%) dbj BAB04138.1  (AP001508) transcriptional regulator (GntR family) [Bacillus halodurans] Length = 240
-Gontig125 (25927-266	 Contig125 (25443-255	Contig125 (25003-254	Contig125 (24286-247	Contig125 (23406-242	Contig131 (43560-45074 m)	Contig125 (22530-232		Contig125 (20306-210
SA-735.1	SA-736.1	SA-737.1	SA-738.1	SA-739.1	SA-74.1	SA-740.1	SA-741.1	SA-742.1
SeqID 2100	SeqID 2101	SeqID 2102	SeqID 2103	SeqID 2104	SeqID 2105	SeqID 2106	SeqID 2107	SeqID 2108



SeqID 2109	SA-743.2	Contig125 (18531-20093 m)	88	Identities = 418/511 (81%), Positives = 467/511 (90%), Gaps = 3/511 (0%) gb AAK05584:1 AE006379_1-(AE006379) GMP synthase (EC 6.3.5.2) [Lactococcus lactis subsp. Length = 513
SeqID 2110	SA-744.2	Contig119 (32996-33796 m)	57	Identities = 113/251 (45%), Positives = 160/251 (63%), Gaps = 3/251 (1%) gb AAK04846.1 AE006308_6 (AE006308) HYPOTHETICAL PROTEIN [Lactococcus lactis subsp. lactis] Length = 257
SeqiD 2111	SA-745.1	Contig119 (32371-33006 m)	74	Identities = 126/211 (59%), Positives = 161/211 (75%) pir  B69997 conserved hypothetical protein ytmQ - Bacillus subtilis gb AAC00285.1  (AF008220) YtmQ {Bacillus subtilis} emb CAB14968.1  (Z99119) similar to hypothetical proteins [Bacillus subtilis]
SeqID 2112	SA-746.1	Contig119 (31405-31899 m)	09	Identities = 73/152 (48%), Positives = 99/152 (65%), Gaps = 3/152 (1%) dbj BAB06136.1  (AP001515) unknown conserved protein [Bacillus halodurans] Length = 156
SeqID 2113	SA-747.1	Contig119 (30218-31369 m)	63	Identities = 164/370 (44%), Positives = 251/370 (67%), Gaps = 15/370 (4%) splP32727 NUSA_BACSU N UTILIZATION SUBSTANCE PROTEIN A HOMOLOG emb CAB13533.1  (299112) nusA [Bacilius subtilis] Length = 371
SeqiD 2114	SA-748.1	Contig119 (29900-30196 m)	65	Identities = 46/92 (50%), Positives = 67/92 (72%), Gaps = 1/92 (1%) splP32728 YLXR_BACSU HYPOTHETICAL 10.4 KD PROTEIN IN NUSA-INFB INTERGENIC REGION (ORF3) pir  D36905 conserved hypothetical protein ylxR - Bacillus subtilis emb CAA79232.1 (Z18631) ORF3 [Bacillus subtilis] emb CAB13534.1 (Z99112) alternate gene name: ymxB~similar to hypothetical proteins [Bacillus subtilis] Length = 91
SeqID 2115	SA-749.1	Contig119 (29605-29907 m)	29	Identities = 50/97 (51%), Positives = 69/97 (70%) sp P55768 YLXQ_ENTFC PROBABLE RIBOSOMAL PROTEIN IN INFB 5 REGION Length = 103
SeqID 2116	SA-75.1	Contig131 (40941-43481 p)	38	itives : D8508 = 610

SeqID 2117	SA-751.1	Contig119 (26802-29585 m)	86	Identities = 926/927 (99%), Positives = 927/927 (99%)  —spl@9ZF20IF2_STRAG TRANSLATION INITIATION FACTOR—  IF-2 emb CAA05919.1  (AJ003164) initiation factor IF2 [Streptococcus agalactiae] emb CAC00489.1  (AJ251495)  —initiation factor 2 [Streptococcus agalactiae] emb CAC00491+1   (AJ251496) initiation factor 2 [Streptococcus agalactiae]  Length = 927
SeqID 2118	SA-752.1	Contig119 (26343-26711 m)	200	Identities = 122/122 (100%), Positives = 122/122 (100%) emb CAA05920.1 (AJ003164) ribosome binding factor A [Streptococcus agalactiae] emb CAC00486.1 (AJ251493) ribosome binding factor A [Streptococcus agalactiae] emb CAC00488.1 (AJ251494) ribosome binding factor A [Streptococcus agalactiae] emb CAC00490.1 (AJ251495) ribosome binding factor A [Streptococcus agalactiae] emb CAC00492.1 (AJ251496) ribosome binding factor A [Streptococcus agalactiae] emb CAC00498.1 (AJ251499) ribosome binding factor A [Streptococcus agalactiae] Length = 122
SeqID 2119	SA-753.1	Contig119 (25234-26259 p)	26	Identities = 58/220 (26%), Positives = 90/220 (40%), Gaps = 8/220 (3%) splQ01109jBAH_STRHY ACETYL-HYDROLASE gblAA79277.1 (M64783) acetyl-hydrolase [Streptomyces hygroscopicus] Length = 299
SeqID 2120	SA-754.1	Contig119 (24675-25091 m)	89	Identities = 67/138 (48%), Positives = 99/138 (71%) gb AAG10085.1 AF296446_1 (AF296446) CopY (Streptococcus mutans)
SeqID 2121	SA-756.1	Contig119 (22428-24662 m)	. 92	Identities = 440/740 (59%), Positives = 571/740 (76%), Gaps = 1/740 (0%) gblAAG10086.1 AF296446_2 (AF296446) CopA [Streptococcus mutans]
SeqID 2122	SA-757.1	Contig119 (22181-22387 m)	28	Identities = 31/67 (46%), Positives = 43/67 (63%) gb AAG10087.1 AF296446_3 (AF296446) CopZ [Streptococcus mutans] Length = 67



SA-758.1	Contig119 (21457-22071 m)	65	Identities = 107/192 (55%), Positives = 137/192 (70%) spl032216 YVGT_BAGSU-HYPOTHETICAL 21.7 KDA PROTEIN IN SSPG-SECG INTERGENIC REGION pir[JA70041 conserved hypothetical protein yvgT - Bacillus subtilis emb[CAB15351.1] (299121) similar to hypothetical proteins [Bacillus subtilis] Length = 202
ડે	Contig119 (20630-21460 m)	99	Identities = 138/238 (57%), Positives = 184/238 (76%) emb CAB94816.1  (AJ245582) hypothetical protein [Streptococcus thermophilus] Length = 240
Ö	Contig131 (40208-40957 p)	44	Identities = 58/255 (22%), Positives = 115/255 (44%), Gaps = 26/255 (10%) ref[NP_069699.1  carboxylesterase (est-1) [Archaeoglobus fulgidus] pir[JA69358 carboxylesterase (est-1) homolog - Archaeoglobus fulgidus gb[AAB90371.1  (AE001044) carboxylesterase (est-1) [Archaeoglobus fulgidus] Length = 266
Con	Contig119 (17875-20517 m)	83	Identities = 655/879 (74%), Positives = 748/879 (84%), Gaps = 4/879 (0%) splP13252IDPO1_STRPN DNA POLYMERASE I (POL I) pir  A32949 DNA-directed DNA polymerase (EC 2.7.7.7) Streptococcus pneumoniae gb AAA26954.1  (J04479) DNA polymerase I (Streptococcus pneumoniae)
So	Contig119 (17405-17845 m)	28	Identities = 72/134 (53%), Positives = 94/134 (69%), Gaps = 3/134 (2%) dbj BAB05860.1  (AP001514) unknown conserved protein [Bacillus halodurans] Length = 137
ဝိ	Contig119 (16844-17323 m)	75	Identities = 113/156 (72%), Positives = 135/156 (86%), Gaps = 1/156 (0%) gb AAB18795.1  (U76538) Fur-like protein [Streptococcus pyogenes]
Š	Contig119 (15126-16691 m)	1	Identities = 41/146 (28%), Positives = 63/146 (43%), Gaps = 13/146 (8%) emb[CAA06650.1  (AJ005645) sdrc [Staphylococcus aureus] Length = 947

-SeqID-2:130-	— SA-766.1	Contig119 (14327-15013 m)-		Identities = 143/228 (62%), Positives = 183/228 (79%), Gaps = 1/228 (0%) emb CAB54578.1  (AJ006397)-response-regulator   Streptococcus pneumoniae] Length = 232
SeqID 2131	SA-767.1	Contig119 (13288-14325 m)	12	Identities = 190/343 (55%), Positives = 249/343 (72%) emb[CAB54579.1  (AJ006397) histidine-Kiñase-[Streptococcus pneumoniae] Length = 350
SeqID 2132	SA-768.2	Contig119 (12486-13274 p)	23	identities = 35/119 (29%), Positives = 62/119 (51%), Gaps = 15/119 (12%) pir  S32215 hypothetical protein 1 - Bacillus megaterium emb  CAA79984.1  (Z21972) ORF1 [Bacillus megaterium]
SeqID 2133	SA-769.2	Contig119 (11205-12347 m)	. 08	identities = 269/377 (71%), Positives = 320/377 (84%) sp[032053]TGT_BACSU QUEUINE TRNA-RIBOSYLTRANSFERASE (TRNA-GUANINE TRANSGLYCOSYLASE) (GUANINE INSERTION ENZYME) pir[]B69722 queuine tRNA-ribosyltransferase (EC 2.4.2.29) - Bacillus subtilis emb[CAB14731.1] (Z99118) tRNAguanine transglycosylase [Bacillus subtilis] (Y15896) tRNA-guanine transglycosylase [Bacillus subtilis] Length = 381
SeqID 2134	SA-77.1	Contig131 (39724-40080 p)	89	Identities = 70/119 (58%), Positives = 89/119 (73%), Gaps = 1/119 (0%) splQ02009 YTRP_LACLA HYPOTHETICAL 13.3 KD PROTEIN IN TRPE 5 REGION pir  S35123 hypothetical protein (trpE 5 region) - Lactococcus lactis subsp. lactis gb AAA25222.1  (M87483) ORF 1 [Lactococcus lactis] Length = 119
SeqID 2135	SA-770.1	Contig119 (10787-11098 m)	59	Identities = 45/98 (45%), Positives = 65/98 (65%), Gaps = 3/98 (3%) pir  T38499 hypothetical protein SPAC29B12.12 - fission yeast (Schizosaccharomyces pombe) emb CAB16256.1  (299164) hypothetical protein [Schizosaccharomyces pombe] Length = 113



ı				Identities = 62/173 (35%), Positives = 97/173 (55%), Gaps =
o,	SA-772.1	Contig119 (10241-10780 m)	53	1/173 (0%) pir  B69832 biotin biosynthesis homolog yhfU - Bacillus subtilis emb CAA74544.1  (Y14084) hypothetical protein [Bacillus subtilis] emb CAB12877.1  (Z99109) similar to biotin - biosynthesis [Bacillus subtilis] Length = 186
	SA-773.1	Contig119 (9326-10102 m)	48	Identities = 89/218 (40%), Positives = 129/218 (58%), Gaps = 4/218 (1%) pir  F69830 conserved hypothetical protein yhfl - Bacillus subtilis emb CAA74530.1  (Y14083) hypothetical protein [Bacillus subtilis] emb CAB12864.1  (299109) similar to hypothetical proteins [Bacillus subtilis] Length = 244
	SA-774.2	Contig119 (8820-9326 m)	<b>2</b> 9	Identities = 71/154 (46%), Positives = 110/154 (71%) sp P21335 YAAJ_BACSU HYPOTHETICAL 17.8 KD PROTEIN IN SERS-DNAH INTERGENIC REGION pir  S11690 conserved hypothetical protein yaaJ - Bacillus subtilis emb CAA36389.1  (X52144) ORF17 (AA 1-161) [Bacillus subtilis] dbj BAA05254.1  (D26185) unknown [Bacillus subtilis] emb CAB11794.1  (Z99104) similar to hypothetical proteins [Bacillus subtilis] prf  1617102A 17kD protein [Bacillus subtilis]
	SA-776.1	Contig110 (3335-3646 p)	No Hits found	
	SA-777.1	Contig110 (2062-2373 m)	27	Identities = 25/35 (71%), Positives = 29/35 (82%) pir  G81516 hypothetical protein CP0988 [imported] - Chlamydophila pneumoniae (strain AR39) Length = 52
1 1	SA-778.1		No Hits found	
- 1	SA-779.1	Contig110 (1371-1676 m)	No Hits found	

SeqID 2143	SA-78.1	Contig131 (38025=39656.p).	8	Identities = 159/568 (27%), Positives = 265/568 (45%), Gaps = 82/568 (14%) splP32399JYHGE_BACSU HYPOTHETICAL 84.1 KD PROTEIN IN HEMY-GLTT INTERGENIC REGION (ORFB) pir  H69832 phage infection protein homolog yhgE - Bacillus subtilis emb CAA74522.1[(Y14083) hypothetical protein [Bacillus subtilis] emb CAB12856.1] (Z99109) alternate gene name: yixE~similar to phage infection protein [Bacillus subtilis]
SeqID 2144	SA-780.1	Contig110 (499-846 m)	No Hits found	
SeqID 2145	SA-782.2	Contig139 (106281-108005 p)	46	Identities = 122/553 (22%), Positives = 265/553 (47%), Gaps = 12/553 (2%) dbj BAB06924.1  (AP001518) BH3205~unknown conserved protein [Bacillus halodurans] Length = 561
SeqID 2146	SA-783.1	Contig139 (108099-108740 p)	64	Identities = 94/212 (44%), Positives = 137/212 (64%) pir  E70860 probable serB2 protein - Mycobacterium tuberculosis (strain H37RV) emb CAA16127.1  (AL021287) serB2 [Mycobacterium tuberculosis] Length = 409
SeqID 2147	SA-784.1	Contig139 (108761-109246 m)	24	Identities = 24/77 (31%), Positives = 40/77 (51%) gb AAG20030.1  (AE005083) isopentenyl pyrophosphate isomerase; Idi [Halobacterium sp. NRC-1] Length = 213
SeqID 2148	SA-785.1	Contig139 (109259-109714 m)	36	Identities = 32/116 (27%), Positives = 57/116 (48%), Gaps = 4/116 (3%) pir  B70008 hypothetical protein yuel - Bacillus subtilis emb CAB15165.1  (299120) yuel [Bacillus subtilis] Length = 132
SeqID 2149	SA-786.1	Contig139 (109912-111219 p)	92	Identities = 396/435 (91%), Positives = 414/435 (95%), Gaps = 1/435 (0%) dbj BAA81815.1  (AB029313) enolase [Streptococcus intermedius]
SeqID 2150	SA-787.1	Contig139 (111327-112391 m)	30	Identities = 62/236 (26%), Positives = 107/236 (45%), Gaps = 38/236 (16%) gb AAF20148.1 AF208390_1 (AF208390) actininlike protein [Entamoeba histolytica] Length = 537



SeqID 2151	SA-788.1	Contig139 (112620-113903 p)	79	Identities = 287/426 (67%). Positives = 346/426 (80%) spiQ9S400jAROA_STRPN 3-PHOSPHOSHIKIMATE 1- CARBOXYVINYLTRANSFERASE (5- ENOLPYRUVYLSHIKIMATE-3-PHOSPHATE SYNTHASE) (EPSP SYNTHASE) (EPSPS) gbjAAD45819 1jAF 169483_1 (AF 169483) 5-enolpyruvylshikimate- 3-phosphate synthase [Streptococcus pneumoniae] Length = 427
SeqID 2152	SA-789.1	Contig139 (113896-114408 p)	99	Identities = 65/164 (39%), Positives = 98/164 (59%), Gaps = 8/164 (4%) sp P43906 AROK_LACLA SHIKIMATE KINASE (SK) pir  S52581 shikimate kinase (EC 2.7.1.71) - Lactococcus lactis emb CAA55181.1  (X78413) shikimate kinase [Lactococcus lactis] Length = 162
SeqID 2153	SA-79.1	Contig131 (37307-38101 p)	51	Identities = 109/229 (47%), Positives = 165/229 (71%) gb AAC14610.1  (U95842) transmembrane protein Tmp7 [Lactococcus lactis] Length = 234
SeqID 2154	SA-790.1	Contig139 (114432-115838 p)	43	Identities = 143/409 (34%), Positives = 206/409 (49%), Gaps = 56/409 (13%) emb CAB76821.1  (AJ276231) PSR protein [Enterococcus faecalis] emb CAB76822.1  (AJ276232) PSR protein [Enterococcus faecalis]
SeqID 2155	SA-792.1	Contig139 (115939-117294 p)	63	Identities = 198/452 (43%), Positives = 300/452 (65%) dbj BAB04406.1  (AP001509) RNA methyltransferase [Bacillus halodurans] Length = 458
SeqID 2156	SA-793.1	Contig139 (117321-117623 p)	No Hits found	
SeqID 2157	SA-794.1	Contig139 (117741-118478 p)	28	Identities = 112/245 (45%), Positives = 148/245 (59%), Gaps = 10/245 (4%) pir[ T09434 acid phosphatase (EC 3.1.3.2) [similarity] - Haemophilus influenzae (strains CCUG, Rd KW20) emb CAA68889.1  (Y07615) acid phosphatase [Haemophilus influenzae] Length = 235

SeqID 2158	SA-795.1	Contig139 (118799-119317 p)	29	Identities = 93/173 (53%), Positives = 123/173 (70%), Gaps = 3/173-(1%)-ref[NP_010829.1  Ydr540cp [Saccharomyces cerevisiae] pir  S62019 hypothetical protein YDR540c - yeast (Saccharomyces cerevisiae) gb AAB64982.1  (U43834) Ydr540cp [Saccharomyces cerevisiae] Length = 179
SeqID 2159	SA-796.1	Contig139 (119446-119634 m)	No Hits found	
SeqID 2160	SA-797.1	Contig139 (119663-120118 m)	32	Identities = 29/95 (30%), Positives = 49/95 (51%), Gaps = 2/95 (2%) gb AAG09969.1 AF248037_4 (AF248037) unknown [Streptococcus agalactiae] Length = 174
SeqID 2161	SA-798.1	Contig139 (120154-120486 p)	34	Identities = 31/63 (49%), Positives = 44/63 (69%), Gaps = 2/63 (3%) gb AAG01392.1 AF208158_1 (AF208158) alpha-like protein 2 [Streptococcus agalactiae] Length = 786
SeqID 2162	SA-799.1	Contig139 (120607-121428 m)	92	Identities = 285/296 (96%), Positives = 287/296 (96%) pir[JT09011 probable transposase TnpA - Streptococcus pyogenes (fragment) gb[AAB92607.1  (AF026542) TnpA [Streptococcus pyogenes] Length = 364
SeqID 2163	SA-8.1	Contig137 (36802-39009 p)	28	Identities = 123/560 (21%), Positives = 212/560 (36%), Gaps = 74/560 (13%) ref NP_053169.1  pXO2-14 [Bacillus anthracis] gb AAF13619.1 AF188935_17 (AF188935) pXO2-14 [Bacillus anthracis] Length = 952
SeqID 2164	SA-80.1	Contig131 (36659-37198 m)	51	Identities = 47/181 (25%), Positives = 95/181 (51%), Gaps = 16/181 (8%) dbj BAB04438.1 (AP001509) transcriptional regulator (TetR/AcrR family) [Bacillus halodurans]
SeqID 2165	SA-800.1	Contig139 (121476-121751 m)	. 61	Identities = 66/80 (82%), Positives = 68/80 (84%) pir  T09011 probable transposase TnpA - Streptococcus pyogenes (fragment) gb AAB92607.1  (AF026542) TnpA (Streptococcus pyogenes] Length = 364



Contig139 (1
Contig139 (122409-122756 m)***
Contig139 (122950-123369 m)
Contig139 (123370-124158 m)
Contig139 (124541-126205 p)
Contig139 (126330-127217 p)
Contig139 (127219-128136 p)
Contig139 (128192-128944 p)
Contig131 (36277-36573 p)

SeqID 2175	SA-810.3	Contig 139 (128806-128955 p)	No Hits found	
SeqID 2176	SA-811.2	Contig 123 (32704-32853 p)	85	Identities = 43/49 (87%). Positives = 46/49 (93%)   spiO34102 RL33_LACLC 50S RIBOSOMAL PROTEIN L33   gb AAB66692 1  (U89998) 50S ribosomal protein subunit L33   Lactococcus lactis subsp. cremoris   Length = 49
SeqID 2177	SA-812.1	Contig123 (32506-32688 p)		Identities = 46/53 (86%), Positives = 49/53 (91%) spjO34101 RL32_LACLC 50S RIBOSOMAL PROTEIN L32 gbjAAB66691.1  (U89998) 50S ribosomal protein subunit L32 [Lactococcus lactis subsp. cremoris] Length = 58
SeqID 2178	SA-814.1	Contig123 (31006-32286 m)	8	Identities = 342/423 (80%), Positives = 377/423 (88%) splP30053 SYH_STREQ HISTIDYL-TRNA SYNTHETASE (HISTIDINETRNA LIGASE) (HISRS) pir  S30233 histidine-tRNA ligase (EC 6.1.1.21) [validated] - Streptococcus equisimilis Length = 426
SeqID 2179	SA-817.1	Contig123 (29159-30913 m)	73	Identities = 339/586 (57%), Positives = 432/586 (72%), Gaps = 10/586 (1%) splO32038 SYD_BACSU ASPARTYL-TRNA SYNTHETASE (ASPARTATETRNA LIGASE) (ASPRS) pir  D69591 aspartatetRNA ligase (EC 6.1.1.12) aspS - Bacillus subtilis emb CAB14714.1  (299118) aspartyl-tRNA synthetase [Bacillus subtilis]
SeqID 2180	SA-819.1	Contig123 (28225-29175 m)	25	Identities = 105/275 (38%), Positives = 182/275 (66%), Gaps = 1/275 (0%) emb CAB01834.1  (Z79580) putative ORF [Bacillus subtilis] emb CAA70630.1  (Y09476) YItT [Bacillus subtilis] emb CAB12952.1  (Z99109) alternate gene name: yuxA~similar to hypothetical proteins [Bacillus subtilis] Length = 280



SA-82.1 Contig131 (35422-36033 p) SA-820.1 Contig123 (27245-28117 m) SA-822.1 Contig123 (26910-27218 p) SA-822.1 Contig123 (25131-26822 p) SA-823.1 Contig123 (24472-24918 m) SA-825.1 Contig123 (21839-24415 m)	77 64	Identities = 138/201 (68%), Positives = 158/201 (77%), Gaps = 1/201 (0%) splP21466IRS4_BACSU 30S RIBOSOMAL PROTEIN S4 (BS4) pir  A37146 ribosomal protein S4 - Bacillus subtilis gb AAA22717.1  (M59358) ribosomal protein S4 [Bacillus subtilis] gb AAA22717.1  (M59358) ribosomal protein S4 [Bacillus subtilis] gb AAC00397.1  (AF008220) ribosomal protein S4 (B34) [Bacillus subtilis] Length = 200 [Bacillus subtilis] Length = 200 Identities = 113/278 (40%), Positives = 192/278 (68%), Gaps = 1/278 (0%) db  BAB05397.1  (AP001512) unknown conserved protein [Bacillus halodurans] Length = 290 Identities = 27/88 (30%), Positives = 48/88 (53%), Gaps = 1/88
	64	Identities = 113/278 (40%), Positives = 192/278 (68%), Gaps = 1/278 (0%) dbj BAB05397.1  (AP001512) unknown conserved protein [Bacillus halodurans] Length = 290 Identities = 27/88 (30%), Positives = 48/88 (53%), Gaps = 1/88
		Identities = 27/88 (30%), Positives = 48/88 (53%), Gaps = 1/88
	43	(1%) dbj BAA11330.1  (D78257) BacB [Enterococcus faecalis] Length = 94
	80	Identities = 377/566 (66%), Positives = 464/566 (81%), Gaps = 5/566 (0%) gb[AAF86984.1 AF282249_3 (AF282249) ArgS [Lactococcus lactis subsp. lactis] Length = 564
	. 58	Identities = 62/139 (44%), Positives = 89/139 (63%) splQ54870JARGR_STRPN ARGININE REPRESSOR pir. B28667 hypothetical protein C - Streptococcus pneumoniae gbJAAA88596.1 (M18729) unknown protein [Streptococcus pneumoniae] Length = 148
	<b>8</b>	Identities = 593/858 (69%), Positives = 698/858 (81%), Gaps = 14/858 (1%) splP10564 HEXA_STRPN DNA MISMATCH REPAIR PROTEIN HEXA pir] C28667 DNA mismatch repair protein hexA - Streptococcus pneumoniae gb AAA88597.1  (M18729) mismatch repair protein [Streptococcus pneumoniae] Length = 844
SA-826.1 Contig123 (21579-21782 p)	. 76	Identities = 48/63 (76%), Positives = 56/63 (88%) emb CAA71254.1  (Y10181) cold shock protein [Lactococcus lactis] emb CAA76695.1  (Y17215) cold shock protein B [Lactococcus lactis] Length = 66

S = B B B B B B B B B B B B B B B B B B	a la s	∥ ₹	85 /eg //eg s	SE III SE			
1%), Gap FCH REF otein hex \88600.1	(58%), Gaps rotein LmrP - ) LmrP integra Length = 408	%), Gaps unction D = 203	Gaps = 8  S66073 5) unkno g [Bacillu	8%), Gaps = HYLADENINI YCOSYLASE ADENINE ADENINE Nine-DNA CAA27472.1 [Escherichia ine DNA nia coli] A glycosylas/Length = 187			
13/657 (8 A MISMA r repair pr e gblAA/ ococcus i	36/401 (5/ ane prote 89779) L	2/201 (59%), G holliday junction Length = 203	5 (52%), TEIN pirl 1  (D2618 99104) ve : 86	122/176 (68%), G NA-3-METHYLAD E-DNA GLYCOS) -METHYLADENII methyladenine-Di coli emblCAA27, a coli] gblAAA24 lase I (tag) [Esch nethyladenine DN [Escherichia coli] idenine DNA glyco (12] Length			
6), Positives = 54 XB_STRPN DN. 33589 mismatch ccus pneumonia ir protein [Strept Length = 649	itives = 2; al membi 1918.1  (X us lactis)	tives = 12 >001511) ans]	es = 45/85 (5 /EG PROTEI \A05279.1  (I 820.1  (2991) Length = 86	tives = 12 COLI DNA DENINE- (DNA-3-N (DNA-3-N CM1 3-m Perichia o therichia o			
38%), Pos HEXB_S1   A33589  coccus p  epair prote	entities = 145/401 (36%), Positives = 236/401 (58%), Gaps 4/401 (0%) pir  S58131 integral membrane protein LmrP - actococcus lactis emb CAA61918.1  (X89779) LmrP integrambrane protein [Lactococcus lactis]	4/401 (0%) pirilissers integral memorane protein LmrP - Lactococcus lactis emb CAA61918.1  (X89779) LmrP integral membrane protein [Lactococcus lactis] Length = 408 Identities = 86/201 (42%), Positives = 122/201 (59%), Gaps = 5/201 (2%) dbj BAB04943.1  (AP001511) holliday junction DN/helicase [Bacillus halodurans] Length = 203 Identities = 24/85 (28%), Positives = 45/85 (52%), Gaps = 8/85 protein - Bacillus subtilis dbj BAA05279.1  (D26185) unknown [Bacillus subtilis] emb CAB11820.1  (299104) veg [Bacillus subtilis] Length = 86		ntities = 87/176 (49%), Positives = 122/176 (68%), Gap: 6 (0%) sp P05100 3MG1_ECOLI DNA-3-METHYLADEN/COSYLASE I (3-METHYLADENINE-DNA GLYCOSYLASE I (3-METHYLADENINE-DNA GLYCOSIDASE I) pir  DGECM1 3-methyladenine-DNA cosylase (EC 3.2.2) I - Escherichia coli emb CAA274723845) TAGI (aa 1-187) [Escherichia coli gb AAA24658 (2606) 3-methyladenine-DNA glycosylase I (tag) [Escherichia coli] gb AAB18526.1 (1000039) 3-methyladenine DNA glycosylase I, constitutive [Escherichia coli] abcosylase I, constitutive [Escherichia coli] constitutive [Escherichia coli] Length = 'Constitutive [Escherichia coli K12]			
152/657 (6 0 P14160  HEXB pii 1] - Streptc ismatch re	145/401 (3 b) pir  S58 s lactis er protein [	membrane protein [Lactococcus lactis] Length = 408 Identities = 86/201 (42%), Positives = 122/201 (59%), Gaps = 6/201 (2%) dbj BAB04943.1  (AP001511) holliday junction DNA helicase [Bacillus halodurans] Length = 203 Identities = 24/85 (28%), Positives = 45/85 (52%), Gaps = 8/85 (9%) splp37466 VEG_BACSU VEG PROTEIN pirl S66073 veg protein - Bacillus subtilis dbj BAA05279.1  (D26185) unknown [Bacillus subtilis] emb CAB11820.1  (Z99104) veg [Bacillus subtilis]		87/176 (4 ppP05100  ASE I (3-h TITUTIVE SIDASE I) (EC 3.2.2 AGI (aa 1 nethylade  AAB1852 sylase I, c			
Identities = 452/657 (68%), Positives = 543/657 (81%), Gaps = 8/657 (1%) sp P14160 HEXB_STRPN DNA MISMATCH REPAIR PROTEIN HEXB pir  A33589 mismatch repair protein hexB [validated] - Streptococcus pneumoniae gb AAA88600.1  (M29686) mismatch repair protein [Streptococcus pneumoniae] Length = 649	[validated] - Streptococcus pneumoniae gb AAA88600.1  [M29686) mismatch repair protein [Streptococcus pneumoniae] Length = 649  Identities = 145/401 (36%), Positives = 236/401 (58%), Gaps = 4/401 (0%) pir  S58131 integral membrane protein LmrP - Lactococcus lactis emb CAA61918.1  (X89779) LmrP integral membrane protein [Lactococcus lactis]  Identities = 86/201 (42%), Positives = 122/201 (59%), Gaps = 6/201 (2%) dbj BAB04943.1  (AP001511) holliday junction DNA helicase [Bacillus halodurans]  Identities = 24/85 (28%), Positives = 45/85 (52%), Gaps = 8/85 (9%) splP37466 VEG_BACSU VEG PROTEIN pir  S66073 veg protein - Bacillus subtilis dbj BAA05279.1  (D26185) unknown [Bacillus subtilis] emb CAB11820.1  (299104) veg [Bacillus subtilis]		Identities = 87/176 (49%), Positives = 122/176 (68%), Gaps = 1/176 (0%) splP05100 3MG1_ECOLI DNA-3-METHYLADENINE GLYCOSYLASE I (3-METHYLADENINE-DNA GLYCOSYLASE I, CONSTITUTIVE) (TAG I) (DNA-3-METHYLADENINE GLYCOSIDASE I) pir  DGECM1 3-methyladenine-DNA glycosylase (EC 3.2.2) I - Escherichia coli emb CAA27472.1  (X03845) TAGI (aa 1-187) [Escherichia coli] gb AAA24658.1  (J02606) 3-methyladenine-DNA glycosylase I (tag) [Escherichia coli] gb AAB18526.1  (U00039) 3-methyladenine DNA glycosylase I, constitutive [Escherichia coli] ablyAB18526.1  (Becherichia coli K12] Length = 187				
9/8 - W	W)   P		P 60	2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
62	55	61	44				
· · · · · · · · · · · · · · · · · · ·	:						
359 m)	348 m)	3348 m) 3114 m) 5092 p)		.501 m)			
19380-21359 m)	18116-19348 m)	17524-18114 m)	(34805-35092 p)	16950-17501 m)			
	Contig 123 (1)		Contig131 (3	Contig123 (16			
Co	Contig123 Contig123 Contig133		Cont				
۸-827.1	SA-827.1 SA-828.1 SA-829.1		SA-831.1				
		·	·				
eqID 2188	SeqID 2188 SeqID 2189		SeqID 2191	SeqID 2192			
S	S	S	S	Ø			



SeqID 2193	SA-832.1	Contig123 (15602-16861 m)	08	Identities = 292/417 (70%), Positives = 343/417 (82%), Gaps = 1/417 (0%) splP54184 CINA_STRPN PUTATIVE COMPETENCE_DAMAGE PROTEIN (EXPORTED PROTEIN 10) emb CAA84071.1  (Z34303) CinA protein [Streptococcus pneumoniae] Length = 418
SeqID 2194	SA-833.1		88	%), Pc 1 (AF3
SeqID 2195	SA-834.1	Contig123 (13775-14185 m)	29	Identities = 61/127 (48%), Positives = 93/127 (73%)   gb AAF21893.1 AF103794_1 (AF103794) unknown [Listeria   monocytogenes]
SeqID 2196	SA-835.1	Contig123 (13307-13573 m)	70	tives = P0015
SeqID 2197	SA-837.1	Contig123 (12888-13298 m)		Identities = 82/138 (59%), Positives = 109/138 (78%), Gaps = 1/138 (0%) spj034634 YRRK_BACSU HYPOTHETICAL 15.2 KDA PROTEIN IN UDK-ALAS INTERGENIC REGION pir  D69979 conserved hypothetical protein yrrK - Bacillus subtilis emb CAB14681.1  (Z99117) similar to hypothetical proteins [Bacillus subtilis] emb CAB14698.1  (Z99118) similar to hypothetical proteins [Bacillus subtilis]
SeqID 2198	SA-838.1	Contig123 (12545-12862 m)	09	Identities = 50/95 (52%), Positives = 66/95 (68%), Gaps = 5/95 (5%) pir  A69982 hypothetical protein yrzB - Bacillus subtilis emb CAB14680.1  (299117) yrzB [Bacillus subtilis] emb CAB14697.1  (299118) yrzB [Bacillus subtilis] Length = 93
SeqID 2199	SA-839.2	Contig123 (10754-12295 m)	4	Identitles = 39/172 (22%), Positives = 75/172 (42%), Gaps = 9/172 (5%) ref[NP_066532.1] ABC transporter subunit [Naegleria gruberi] gb[AAG17810.1]AF288092_35 (AF288092) ABC transporter subunit [Naegleria gruberi]

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Identities = 398/452 (88%), Positives = 432/452 (95%), Gaps :	delta prime subunit [Streptococcus pyogenes]  Length = 455	Identities = 193/426 (45%), Positives = 278/426 (64%), Gaps = 28/426 (6%) pir  B82263 PTS system, trehalose-specific IIBC component VC0910 [imported] - Vibrio cholerae (group O1 strain N16961) gb AAF94072.1  (AE004175) PTS system, trehalose-specific IIBC component [Vibrio cholerae] Length = 478	Identities = 380/547 (69%), Positives = 437/547 (79%), Gaps   13/547 (2%) gb AAB65079.1  (U35633) dextran glucosidase   DexS [Streptococcus suis]	Identities = 203/697 (29%), Positives = 362/697 (51%), Gaps = 34/697 (4%) dbj BAB03939.1  (AP001507) unknown conserved protein [Bacillus halodurans]	Identities = 26/82 (31%), Positives = 52/82 (62%), Gaps = 2/82 (2%) dbj BAB03941.1  (AP001507) unknown conserved protein [Bacillus halodurans]	Identities = 203/453 (44%), Positives = 293/453 (63%), Gaps = 22/453 (4%) pir T37066 probable integral membrane protein - Streptomyces coelicolor emb CAB52363.1  (AL109747) putative integral membrane protein [Streptomyces coelicolor A3(2)] Length = 516		Identities = 105/269 (39%), Positives = 158/269 (58%), Gaps = 4/269 (1%) splQ58094 TKTN_METJA PUTATIVE TRANSKETOLASE N-TERMINAL SECTION (TK) pir  A64385 transketolase (EC 2.2.1.1) - Methanococcus jannaschii gb AAB98676.1  (U67515) transketolase [Methanococcus jannaschii] Length = 274
	96	45	80	53	99	99	No Hits found	54
	Contig131 (33438-34793 p)	Contig112 (127-1845.p)	Contig112 (2067-3692 p)	Contig112 (3912-5948 p)	Contig112 (5951-6235 p)	Contig112 (6248-7603 p)	Contig112 (7542-7700 m)	Contig112 (7606-8463 p)
	SA-84.1	SA-842.2	SA-843.1	SA-844.1	SA-845.1	SA-846.1	SA-847.1	SA-848.1
	SeqID 2200	SeqID 2201	SeqID 2202	SeqID 2203	SeqID 2204	SeqID 2205	SeqID 2206	SeqID 2207



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(54%), Gal UTATIVE – IK) pir  G6 Is jannasch ethanococc	105/149 (69%), Gaps = SRIBOSOMAL PROTE!! protein L9 - Bacillus in K, Seven Ribosomal Microscopic Map Of Angstroms Resolution Length = 149	5/438 (44%), Ga ein PA0545 [imp (strain PAO1) ) hypothetical pro Length = 434	71/89 (78%) IAL PROTEIN \$ 5 (rpsO) - Bacil somal protein S 2) ribosomal pr Length = 89	
Identities = 99/310 (31%), Positives = 172/310 (54%), Gaps = TRANSKETOLASE C-TERMINAL SECTION (TK) pir  G64384 transketolase (EC 2.2.1.1) - Methanococcus jannaschii gb AAB98674.1  (U67515) transketolase [Methanococcus _ jannaschii] Length = 316	Identities = 80/149 (53%), Positives = 105/149 (69%), Gaps = 149 (1%) splP02417IRL9_BACST 50S RIBOSOMAL PROTEIL	Identities = 111/438 (25%), Positives = 195/438 (44%), Gaps = 1/438 (11%) pir][G83576 hypothetical protein PA0545 [importec - Pseudomonas aeruginosa (strain PAO1) gb AAG03934 1 AE004491_1 (AE004491) hypothetical protein [Pseudomonas aeruginosa] Length = 434	Identities = 55/89 (61%), Positives = 71/89 (78%) splP21473 RS15_BACSU 30S RIBOSOMAL PROTEIN S15 (BS18) pirl F69700 ribosomal protein S15 (rpsO) - Bacillus subtilis emb CAB02560.1  (Z80835) ribosomal protein S15 acillus subtilis] emb CAB13541.1  (Z99112) ribosomal protein S15 (BS18) [Bacillus subtilis]	
%), Positive 58092 TKTC FERMINAL 2.2.1.1) - Me 515) transk	Identities = 80/149 (53%), Positives = 149 (1%) splP02417IRL9_BACST 508 L9 (BL17) pirIIR5BS7F ribosomal stearothermophilus pdbl487DIK Cha Proteins Fitted To A Cryo-Electron The Large 50s Subunit At 7.5 pdbl1DIV  Ribosomal Protein L9	es = 111/438 (25%), Positive 11%) pir  G83576 hypothetic - Pseudomonas aeruginosa 303934. 1 AE004491_1 (AE0 [Pseudomonas aeruginosa]	dentities = 55/89 (61%), Posi 1473 RS15_BACSU 30S RIB 3) pir  F69700 ribosomal prot is emb CAB02560.1  (Z8083 is subtilis] emb CAB13541.1  S15 (BS18) [Bacilius subtilis]	
99/310 (31%), 0 (2%) splQ580 OLASE C-TER tolase (EC 2.2.1674.1  (U67519 jannaschil]	Jentities = 80/149 (53%), Posi 49 (1%) spIPO2417IRL9_BAC L9 (BL17) pirIIR5BS7F ribo stearothermophilus pdbI487D roteins Fitted To A Cryo-Elect The Large 50s Subunit At 7.5 pdbJ1DIV  Ribosomal Prot	111/438 (25) pir  G8357 eudomonas 34.1 AE004 eudomonas	ities = 55/8( IRS15_BA( Ir F69700 ri Ir CAB025 IRI(IRI) emb C	
Identities = 7/310 TRANSKET transke gb AAB96	Identities = 80/149 (53%), Positives = 105/149 (69%), Gaps = 2/149 (1%) splP02417 [RL9_BACST 50S RIBOSOMAL PROTEIN L9 (BL17) pir[ R5BS7F ribosomal protein L9 - Bacillus stearothermophilus pdb 487D [K Chain K, Seven Ribosomal Proteins Fitted To A Cryo-Electron Microscopic Map Of The Large 50s Subunit At 7.5 Angstroms Resolution pdb 1D [V] Ribosomal Protein L9 Length = 149	Identities = 111/438 (25%), Positives = 195/438 (44%), Gaps = 51/438 (11%) pir  G83576 hypothetical protein PA0545 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG03934.1 AE004491_1 (AE004491) hypothetical protein [Pseudomonas aeruginosa] Length = 434	Identities = 55/89 (61%), Positives = 71/89 (78%) splP21473 RS15_BACSU 30S RIBOSOMAL PROTEIN S15 (BS18) pir  F69700 ribosomal protein S15 (rpsO) - Bacillus subtilis emb CAB02560.1  (Z80835) ribosomal protein S15 (Bacillus subtilis] emb CAB13541.1  (Z99112) ribosomal protein S15 (BS18) [Bacillus subtilis] Length = 89	
			0	
90	67	42	70	
389 p)	3395 p)	757 p)	114 p)	
2 (8460-9389 p)	Contig131 (32943-33395 p)	Contig112 (9498-10757 p)	(10845-11114 p)	
Contig112	Contig131	Contig112 (9		
SA-849.1	SA-85.1	SA-850.1	SA-851.1	
SeqID 2208	SeqID 2209	SeqiD 2210	SeqID 2211	
Seq	Seq	Seq	Seq	

SeqID 2212	SA-852.1	Contig112 (11495-13624 p)	47	Identities = 431/708 (60%), Positives = 538/708 (75%), Gaps = 8/708 (1%) splP50849 PNPA_BACSU POLYRIBONUCLEOTIDE NUCLEOTIDYLTRANSFERASE (POLYNUCLEOTIDE PHOSPHORYLASE) (VEGTATIVE PROTEIN 15) (VEG15) pir  S70691 polyribonucleotide nucleotidyltransferase (EC 2.7.7.8) alpha chain pnpA - Bacillus subtilis gb AAC43595.1  (U29668) polynucleotide phosphorylase (Bacillus subtilis] emb CAB13542.1  (Z99112) polynucleotide phosphorylase (Bacillus subtilis] Length = 705
-	SA-853.1	Contig112 (13626-14378 p)	No Hits found	
	SA-854.1	Contig112 (14387-14971 p)	. 19	Identities = 92/169 (54%), Positives = 125/169 (73%) ·emb CAB71304.1  (AJ130879) serine acetyltransferase [Clostridium sticklandii] Length = 191
LJ	SA-855.1	Contig112 (14972-15163 p)	No Hits found	
	SA-856.1	Contig112 (15232-16503 p)		Identities = 246/467 (52%), Positives = 323/467 (68%), Gaps = 23/467 (4%) splQ06752 SYC_BACSU CYSTEINYL-TRNA SYNTHETASE (CYSTEINETRNA LIGASE) (CYSRS) pir  C53402 cysteinetRNA ligase (EC 6.1.1.16) cysS - Bacillus subtilis gb AAA21798.1  (L14580) cysteinyl-tRNA synthetase [Bacillus subtilis] dbj BAA05328.1  (D26185) cysteinyl-tRNA synthetase [Bacillus subtilis] emb CAA52167.1  (X73989) cysteinetRNA ligase [Bacillus subtilis] emb CAB11870.1  (Z99104) cysteinyl-tRNA synthetase [Bacillus subtilis] Length = 466
	SA-857.1	Contig112 (16496-16882 p)	. 99	Identities = 58/122 (47%), Positives = 87/122 (70%) pir  C69742 conserved hypothetical protein yazC - Bacillus subtilis emb CAB11871.1  (299104) similar to hypothetical proteins [Bacillus subtilis] Length = 143



Contig112 (16985-17740 p)
Contig112 (17737-18255 p)
Contig131 (31063-32937 p)
Contig112 (18348-19208 p)
Contig112 (20193-21194 m)
Contig127 (33995-35254 p)

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SeqiD 2224	SA-866.2		75	Identities = 445/769 (57%), Positives = 582/769 (74%), Gaps = -9/769 (14%)-emb[CAA05302.1  (AJ002291) penicillin-binding protein 1b [Streptococcus pneumoniae] gb[AAF04736.1 AF101781_1 (AF101781) penicillin-binding protein 1b.[Streptococcus pneumoniae]
	SA-867.1	Contig127 (31366-31455 m)	No Hits found	
	SA-868.1	Contig127 (27488-31063 m)	81	Identities = 814/1173 (69%), Positives = 978/1173 (82%), Gaps = 17/1173 (1%) emb CAB56706.1] (Y16468) DNA-dependent RNA polymerase subunit beta [Listeria monocytogenes] Length = 1184
	SA-87.1	Contig131 (28966-30888 p)	78	Identities = 402/627 (64%), Positives = 503/627 (80%), Gaps = 5/627 (0%) dbj BAB07780.1  (AP001520) glucose-inhibited division protein [Bacillus halodurans] Length = 632
l	SA-872.1	Contig127 (23721-27371 m)	78	Identities = 935/990 (94%), Positives = 959/990 (96%), Gaps = 1/990 (0%) splP95816 RPOC_STRPY DNA-DIRECTED RNA POLYMERASE BETA CHAIN (TRANSCRIPTASE BETA CHAIN) (RNA POLYMERASE BETA SUBUNIT) emb CAA65249.1  (X96385) DNA-dependent RNA polymerase (Streptococcus pyogenes) Length = 989
	SA-874.1	Contig127 (23242-23607 m)	59	Identities = 42/99 (42%), Positives = 75/99 (75%) gb/AAC45309.1  (U81957) putative DNA binding protein [Streptococcus gordonii] Length = 122
	SA-875.1	Contig127 (22098-23069 m)	74	Identities = 202/319 (63%), Positives = 254/319 (79%), Gaps = 1/319 (0%) gb AAC45310.1  (U81957) putative ABC transporter subunit ComYA [Streptococcus gordonii] Length = 319
	SA-876.1	Contig127 (21161-22252 m)	69	Identities = 161/280 (57%), Positives = 218/280 (77%) gb AAC45311.1  (U81957) putative ABC transporter subunit ComYB [Streptococcus gordonii] Length = 282
L	SA-877.1	Contig127 (20835-21164 m)	58	Identities = 63/103 (61%), Positives = 85/103 (82%) gb AAC45312.1  (U81957) ComYC [Streptococcus gordonii] Length = 105



SeqID 2233	SA-878.1	Contig127 (20447-20860 m)	57	Identities = 54/127 (42%), Positives = 83/127 (64%) gb AAC23740 1  (AF052207) competence protein [Streptococcus pneumoniae] Length = 134
SeqID 2234	SA-879.1	Contig127 (20176-20421 m)	49	Identities = 27/91 (29%), Positives = 50/91 (54%) emb[CAA75316 1] (Y15043) hypothetical protein [Lactococcus lactis subspicemoris] Length = 128
SeqID 2235	SA-880.1	Contig127 (19761-20222 m)	99	Identities # 61/147 (41%). Positives = 98/147 (66%), Gaps = 3/147 (2%) emb[CAA75315.1] (Y15043) homology to ComYD from Streptcoccus gordonii, and ComGD from Bacilius subtilis [Lactococcus lactis subsp. cremoris] Length = 150
SeqID 2236	SA-881.1	Contig127 (19412-19783 m)	No Hits found	
SeqID 2237	SA-882.1	Contig127 (18323-19297 m)	53	Identities = 105/329 (31%), Positives = 177/329 (52%), Gaps = 17/329 (5%), SplP37876jYTXK_BACSU HYPOTHETICAL 37.4 KD PROTEIN IN ACKA-SSPA INTERGENIC REGION pirljG70003 hypothetical protein ytkk - Bacillus subtilis gblAAC00317.1 (AF008220) Ytkk [Bacillus subtilis] emblCAB14926.1 (299119) alternate gene name: yth [Bacillus subtilis] Length = 329
SeqID 2238	SA-883.1	Contig127 (17098-18291 m)	73	Identities = 223/395 (56%), Positives = 293/395 (73%), Gaps = 3/395 (0%) splP37877IACKA_BACSU ACETATE KINASE (ACETOKINASE) pirIB49935 acetate kinase (EC 2.7.2.1) ackA-Bacillus subtilis gblAAC36857.1 (L17320) acetate kinase [Bacillus subtilis] emblCAB14925.1 (AF008220) acetate kinase [Bacillus subtilis] emblCAB14925.1 (AF008220) acetate kinase [Bacillus subtilis]
SeqID 2239	SA-884.1	Contig127 (16741-16947 m)	20	Identities = 39/64 (60%), Positives = 49/64 (75%) pir  G75183 probable repressor protein PAB7155 - Pyrococcus abyssi (strain Orsay) emb CAB49550.1  (AJ248284) repressor protein, putative [Pyrococcus abyssi] Length = 73
SeqID 2240	SA-885.2	-	No Hits found	
SeqID 2241	SA-887.2	Contig 139 (52447-52644 p)	No Hits found	

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				Identities = 227/311 (72%), Positives = 269/311 (85%)
SeqID 2242	SA-889.1	Contig139 (52688-53620 m)	82	DEHYDROGENASE (DIHYDROOROTATE OXIDASE) (DHODEHASE) (DHODASE) (DHOD) emb CAB51330.1  (AJ131985) dihydroorotate dehydrogenase [Streptococcus pneumoniae] Length = 311
SeqiD 2243	SA-89.1	Contig131 (28206-28817 p)	. 25	Identities = 71/202 (35%), Positives = 120/202 (59%), Gaps = 5/202 (2%) spj032244 YVBG_BACSU HYPOTHETICAL 22.6 KD PROTEIN IN OPUCA-ENO INTERGENIC REGION pir. F70029 conserved hypothetical protein yvbG - Bacillus subtilis emb CAB15390.1  (299121) similar to hypothetical proteins [Bacillus subtilis]
SeqID 2244	SA-890.1	Contig139 (53807-55042 m)	72	Identities = 237/410 (57%), Positives = 304/410 (73%) emb CAB89121.1  (AJ277485) beta-lactam resistance factor [Streptococcus pneumoniae] Length = 410
SeqID 2245	SA-891.1	Contig139 (55061-56272 m)	02	Identities = 204/410 (49%), Positives = 286/410 (69%), Gaps = 17/410 (4%) emb CAB89537.1  (AJ250766) MurM protein [Streptococcus pneumoniae] emb CAB89539.1  (AJ250767) MurM protein [Streptococcus pneumoniae] Length = 406
SeqID 2246	SA-892.1	Contig139 (56285-57505 m)	09	Identities = 166/410 (40%), Positives = 250/410 (60%), Gaps = 10/410 (2%) emb CAB89120.1  (AJ277484) beta-lactam resistance factor [Streptococcus pneumoniae] Length = 406
SeqID 2247	SA-893.1	Contig139 (57505-58329 m)	<b>89</b>	Identities = 127/269 (47%), Positives = 189/269 (70%), Gaps = 1/269 (0%) splP09997 YIDA_ECOLI HYPOTHETICAL 29.7 KD PROTEIN IN IBPA-GYRB INTERGENIC REGION pir  QQECGB hypothetical 29.7K protein, ibpA-gyrB intergenic region - Escherichia coli (strain K-12) gblAAC76720.1  (AE000446) orf, hypothetical protein [Escherichia coli K12] Length = 270
SeqID 2248	SA-894.1	Contig139 (58388-59704 m)	61	Identities = 212/429 (49%), Positives = 287/429 (66%), Gaps = 5/429 (1%) dbj BAB07537.1  (AP001520) unknown conserved protein [Bacillus halodurans] Length = 432



7	olog PacL ar to	s =	.s = 111) -	Gaps = yhbA - 148.5 kd ternate [Bacillus		ss = ethyl- ins]
	Identities =-378/886-(42%); Positives = 542/886 (60%), Gaps = 45/886 (5%) pirl H69877 calclum-transporting ATPase homolog yloB - Bacillus subtilis emb CAA74269.1  (Y13937) putative PacL protein [Bacillus.subtilis] emb CAB13439.1  (Z99112) similar to calcium-transporting ATPase [Bacillus subtilis]	Identities = 60/221 (27%), Positives = 100/221 (45%), Gaps = 37/221 (16%) pir  E75186 hypothetical protein PAB0011 - Pyrococcus abyssi (strain Orsay) emb CAB48940.1  (AJ248283) hypothetical protein [Pyrococcus abyssi]	Identities = 315/644 (48%), Positives = 447/644 (68%), Gaps = 7/644 (1%) pir  C69621 fructose-bisphosphatase (EC 3.1.3.11) - Bacillus subtilis dbj BAA11277.1  (078193) yydE [Bacillus subtilis] emb CAB16056.1  (299124) fructose-1,6-bisphosphatase [Bacillus subtilis]	Identities = 177/371 (47%), Positives = 242/371 (64%), Gaps = 6/371 (1%) pir  E69820 conserved hypothetical protein yhbA - Bacillus subtilis emb CAB07527.1  (293102) hypothetical 48.5 kd protein [Bacillus subtilis] emb CAB12719.1  (299108) alternate gene name: ygaP~similar to hypothetical proteins   Bacillus subtilis]		dentities = 250/359 (69%), Positives = 292/359 (80%), Gaps = 6/359 (1%) dbj BAB04980:1  (AP001511) (5-methylaminomethyl-2-thiouridylate)-methyltran sferase [Bacillus halodurans]
	Identitie 45/886 ( yloB - Ba protein [ calcium	Identitie 37/22 Pyrococo hypoth	Identitie 7/644 (1* Bacilli su bis	Identitie 6/371 ( Bacillus s protein [ gene nan		Identities 6/359 (19 2-thiouri
No Hits found	22	33	69	64	No Hits found	72
Contig139 (59768-60166 p)	Contig139 (60534-63194 p)	Contig139 (63239-64099 ന)	Contig139 (64251-66182 p)	Contig139 (66272-67396 p)	Contig137 (36200-36682 p)	Contig131 (27053-28174 p)
SA-895.1	SA-896.1	SA-897.1 C	SA-898.1	SA-899.1	SA-9.1 (	SA-90.1
SeqID 2249	SeqID 2250	SeqID 2251	SeqID 2252	SeqID 2253	SeqID 2254	SeqID 2255

SeqID 2256	SA-901.1	Contig139 (67583-68563 p)		Identities = 197/344 (57%), Positives = 255/344 (73%), Gaps = 2/344 (0%) splP28367/RF2_BACSU PEPTIDE CHAIN RELEASE. FACTOR 2 (RF-2) prijJN0146 translation releasing factor RF-2 - Bacillus subtilis gbjAAC673534 tj (AF013188) release factor 2 (Bacillus subtilis) gbjAAC67303 1j (AF017113) putative peptide chain release factor RF-2 (Bacillus subtilis) embjCAB15546 tj (299122) peptide chain release factor 2 (Bacillus subtilis)
SeqID 2257	SA-902.1	Contig139 (68582-69274 p)	76	Identities = 138/228 (60%), Positives = 179/228 (77%) pir  D69627 cell-division ATP-binding protein ftsE - Bacillus subtilis gb AAC67262.1  (AF017113) cell division ATP-binding protein [Bacillus subtilis] emb CAB15543.1  (Z99122) cell-division ATP-binding protein [Bacillus subtilis]
SeqID 2258	SA-904.1	Contig139 (69258-70187 p)	52	Identities = 111/311 (35%), Positives = 181/311 (57%), Gaps = 31/311 (9%) spl034876 FTSX_BACSU CELL DIVISION PROTEIN FTSX HOMOLOG pir  G69627 cell-division protein flsX - Bacillus subtilis gb AAC67264.1  (AF017113) cell division protein [Bacillus subtilis] emb CAB15542.1  (Z99122) cell-division protein [Bacillus subtilis]
SeqiD 2259	SA-905.1	Contig139 (70240-70950 m)	41	Identities = 65/185 (35%), Positives = 100/185 (53%), Gaps = 3/185 (1%) pir][A75409 carboxymethylenebutenolidase-related protein - Deinococcus radiodurans (strain R1) gb]AAF10898.1[AE001979_4 (AE001979) carboxymethylenebutenolidase-related protein [Deinococcus radiodurans] Length = 292
SeqID 2260	SA-906.2	Contig139 (70947-71582 m)	48	Identities = 83/197 (42%), Positives = 114/197 (57%), Gaps = 4/197 (2%) dbj BAB06539.1  (AP001516) BH2820~unknown conserved protein [Bacillus halodurans] Length = 211



SeqID 2261	SA-907:2-	Gontig130 (21596-21796 p)	- 9/	Identities = 56/66 (84%), Positives = 60/66 (90%) -dbj BAA23749.1  (AB009314) proton-translocating ATPase, csubunit [Streptococcus bovis] Length = 67
SeqID 2262	SA-908.2	 Contig130 (21829-22545 p)	75	dbj BAA23750.1  (AB009314) proton-translocating ATPase, a subunit [Streptococcus bovis] Length = 239
SeqID 2263	SA-909.1	Contig130 (22563-23060 p)	73	Identities = 103/165 (62%), Positives = 130/165 (78%) splP95785JATPF_STRMU ATP SYNTHASE B CHAIN pir]JC5737 membrane-bound proton-translocating ATPase (EC 3.6.1) b chain - Streptococcus mutans gbjAAD13379.1  (U31170) ATPase, b subunit [Streptococcus mutans] Length = 165
SeqID 2264	SA-91.1	Contig131 (26139-26807 m)	64	Identities = 101/216 (46%), Positives = 156/216 (71%), Gaps = 2/216 (0%) dbj BAB06216.1  (AP001515) L-serine dehydratase beta subunit [Bacillus halodurans]
SeqID 2265	SA-910.1	Contig130 (23060-23596 p)	89	Identities = 98/178 (55%), Positives = 127/178 (71%) splO50156 ATPD_STRBO ATP SYNTHASE DELTA CHAIN dbj BAA23752.1  (AB009314) proton-translocating ATPase, delta subunit [Streptococcus bovis] Length = 178
SeqID 2266	SA-911.1	Contig130 (23612-25117 p)	86	Identities = 480/501 (95%), Positives = 497/501 (98%) dbj BAA23753.1  (AB009314) proton-translocating ATPase, alpha subunit [Streptococcus bovis] Length = 501
SeqID 2267	SA-912.1	Contig130 (25133-26014 p)	693	Identities = 254/293 (86%), Positives = 278/293 (94%), Gaps = 2/293 (0%) dbj BAA23754.1  (AB009314) proton-translocating ATPase, gamma subunit [Streptococcus bovis] Length = 291
SeqID 2268	SA-913.1	Contig130 (26088-27494 p)	85	Identities = 434/466 (93%), Positives = 454/466 (97%) splP95789 ATPB_STRMU ATP SYNTHASE BETA CHAIN gb AAD13383.1  (U31170) ATPase, beta subunit [Streptococcus mutans] Length = 468

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Identities = 102/138 (73%), Positives = 121/138 (86%), Gaps = 1/138 (0%) dizj BAA23756.1  (AB009314) proton-translocating ATPase, epsiron subunit   Streptococcus bovis   Length = 138	Identities = 248/423 (58%), Positives = 323/423 (75%), Gaps = 5/423 (1%) splP70965 MUA1_BACSU UDP-N-ACETYLGLUCOSAMINE 1-CARBOXYVINYLTRANSFERASE 1 (ENOYLPYRUVATE TRANSFERASE) (UDP-N-ACETYLGLUCOSAMINE ENOLPYRUVYL  TRANSFERASE) (EPT) pir  A69662 UDP-N-acetylglucosamine 1 carboxyvinyltransferase murA - Bacillus subtilis emb CAB03688 1  (Z81356) UDP-N-acetylglucosamine 1 carboxyvinyltransferase [Bacillus subtilis] emb CAB15693.1  (Z99122) UDP-N-acetylglucosamine 1 carboxyvinyltransferase [Bacillus subtilis] Length = 436	Identities = 167/287 (58%), Positives = 200/287 (69%), Gaps = 15/287 (5%) sp Q03158 NUCE_STRPN DNA-ENTRY NUCLEASE (COMPETENCE-SPECIFIC NUCLEASE) pir  S10641 endA protein - Streptococcus pneumoniae emb CAA38134.1  (X54225) membrane nuclease [Streptococcus pneumoniae] Length = 274	Identities = 211/341 (61%), Positives = 272/341 (78%), Gaps = 2/341 (0%) splP17921 SYFA_BACSU PHENYLALANYL-TRNA SYNTHETASE ALPHA CHAIN (PHENYLALANINETRNA LIGASE ALPHA CHAIN (PHERS) pir JYFBSA phenylalaninetRNA ligase (EC 6.1.1.20) alpha chain - Bacillus subtilis emb CAA99603.1  (275208) phenylalany-tRNA synthetase beta subunit [Bacillus subtilis] Length = 344
Contig130-(27507-27920-p)	Contig130 (28278-29549 p)	Contig130 (29817-30674 p)	Contig130 (30965-32005 p)
SA-914.1	SA-916.1	SA-917.1	SA-918.1
SeqID 2269	SeqID 2270	SeqID 2271	SeqiD 2272



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Identities = 61/169 (36%), Positives = 100/169 (59%)  SPORULATION NEGATIVE REGULATORY PROTEIN PAI 1 emb[CAB15205.1] (299120) transcriptional regulator [Bacillus subtilis]  Length = 172	Identities = 376/805 (46%), Positives = 523/805 (64%), Gaps = 6/805 (0%) splP17922 SYFB_BACSU PHENYLALANYL-TRNA SYNTHETASE BETA CHAIN (PHENYLALANINETRNA LIGASE BETA CHAIN) (PHERS) pir JYFBSB phenylalaninetRNA ligase (EC 6.1.1.20) beta chain - Bacillus subtilis emb CAA99564.1  (Z75208) phenylalanyl-tRNA synthetase beta subunit (Bacillus subtilis) emb CAB14823.1  (Z99118) phenylalanyl-tRNA synthetase (beta subunit) [Bacillus subtilis]	Identities = 105/196 (53%), Positives = 141/196 (71%), Gaps = 1/196 (0%) pir  B65023 hypothetical protein b2475 - Escherichia coli (strain K-12) gb AAC75528.1 (AE000334) orf, hypothetical protein [Escherichia coli K12] Length = 287	Identities = 363/1093 (33%), Positives = 606/1093 (55%), Gaps = 67/1093 (6%) pir  T30307 rexB protein - Lactococcus lactis gb AAC12965.1  (U76424) exonuclease RexB [Lactococcus lactis] Length = 1099	Identities = 518/1212 (42%); Positives = 744/1212 (60%), Gaps = 73/1212 (6%) pir  T30308 rexA protein - Lactococcus lactis gb AAC12966.1  (U76424) exonuclease RexA [Lactococcus lactis] Length = 1173	Identities = 88/306 (28%), Positives = 149/306 (47%), Gaps = 17/306 (5%) ref(NP_053049.1  hypothetical protein [Plasmid pNZ4000] gb/AAD40365.1  (AF036485) hypothetical protein [Plasmid pNZ4000]
- 51	64	63	99	61	47
Contig130 (32088-32609 p)	Contig130 (32663-35068 p)	Contig130 (35137-35805 m)	Contig130 (35916-39149 p)	Contig130 (39136-42762 p)	Contig130 (42775-43701 p)
SA-919.1	SA-920.1	SA-921.1	SA-923.1	SA-925.1	SA-926.1
SeqID 2273	SeqiD 2274	SeqID 2275	SeqID 2276	SeqID 2277	SeqID 2278

Identities = 443/462 (95%), Positives = 447/462 (95%), Gaps = 44/462 (0%)-emb[CAB61255.1  (AJ251564) thiophene degradation protein F [Streptococcus agalactiae] Length = 462	Identities = 176/289 (60%), Positives = 224/289 (76%), Gaps = 1/289 (0%) splO34607 SDHA_BACSU PROBABLE L-SERINE DEHYDRATASE, ALPHA CHAIN (L-SERINE DEAMINASE) (SDH) (L-SD) pir  G69879 L-serine dehydratase homolog ylpA - Bacillus subtilis emb CAA74259.1  (Y13937) putative YhaP protein [Bacillus subtilis] emb CAB13459.1  (Z99112) similar to L-serine dehydratase [Bacillus subtilis]	Identities = 241/316 (76%), Positives = 265/316 (83%)     gb AAK05072.1 AE006332_1 (AE006332) ribonucleoside-   diphosphate reductase beta chain (EC	Identities = 367/701 (52%), Positives = 488/701 (69%), Gaps = 19/701 (2%) gb AAD41036.1 AF112535_3 (AF112535)   ribonucleotide reductase alpha-chain [Corynebacterium glutamicum]   Length = 707	Identities = 42/70 (60%), Positives = 53/70 (75%) spiQ48708jNRDH_LACLC GLUTAREDOXIN-LIKE PROTEIN NRDH embjCAA63372.1  (X92690) glutaredoxin-like protein [Lactococcus lactis] Length = 72	Identities = 86/87 (98%), Positives = 87/87 (99%)  dbj BAA78048.1  (AB027569) histidine containing protein  [Streptococcus bovis] Length = 87	Identities = 533/577 (92%), Positives = 559/577 (96%)  dbj BAA78049.1  (AB027569) phosphoenolpyruvate-protein phosphotransferase [Streptococcus bovis] Length = 577
98	92	76	29	67	92	95
Contig130 (43676-45052 m)	Contig131 (25252-26124 m)	Contig117 (7623-8582 m)	Contig117 (8785-10944 m)	Contig117 (11022-11246 m)	Contig117 (11628-11891 p)	Contig117 (11896-13629 p)
SA-928.2	SA-93.1	SA-930.2	SA-931.2	SA-932.2	SA-934.1	SA-936.1

SeqID 2281

SeqID:2283

**SeqID 2282** 

SeqID 2285

SeqID 2284

SeqID 2279

**SeqID 2280** 



	Contig117 (13779-15206 p)	88	SPIQ59931 GAPN_STRMU NADP-DEPENDENT  SPIQ59931 GAPN_STRMU NADP-DEPENDENT  GLYCERALDEHYDE-3-PHOSPHATE-DEHYDROGENASE  (NON-PHOSPHORYLATING GLYCERALDEHYDE 3-PHOSPHATE  3-PHOSPHATE  BEHYDROGENASE) (GLYCERALDEHYDE  3-PHOSPHATE  BEHYDROGENASE) pir JA57151  glyceraldehyde-3-phosphate dehydrogenase (NADP+))  (TRIOSEPHOSPHATE DEHYDROGENASE) pir JA57151  glyceraldehyde-3-phosphate dehydrogenase (NADP+) (EC  1.2.1.9) - Streptococcus mutans pdb 1EUH A Chain A, Apo Form Of A Nadp Dependent Aldehyde Dehydrogenase From Streptococcus Mutans pdb 1EUH C Chain C, Apo Form Of A  Nadp Dependent Aldehyde Dehydrogenase From Streptococcus Mutans pdb 1EUH C Chain C, Apo Form Of A  Nadp Dependent Aldehyde Dehydrogenase From Streptococcus Mutans pdb 1EUH C Chain A, Holo Form Of A  Nadp Dependent Aldehyde Dehydrogenase  Complex With Nadp Dependent Aldehyde Dehydrogenase  Complex With Nadp Dependent Aldehyde Dehydrogenase  Complex With Nadp+ Aldehyde Dehydrogenase  Complex With Nadp+
Conti	Contig117 (15346-16599 p)	54	Identities = 133/439 (30%), Positives = 232/439 (52%), Gaps = 20/439 (4%) emb CAB96552.1  (AJ251472) peptidoglycan GlcNAc deacetylase [Streptococcus pneumoniae]
Contig131	3131 (24440-25117 p)	32	Identities = 70/115 (60%), Positives = 78/115 (66%), Gaps = 6/115 (5%) dbj BAA95959.1  (AB042839) secretory protein SAI-B [Staphylococcus aureus]
Contig	Contig117 (16630-17712 m)	54	Identities = 138/343 (40%), Positives = 200/343 (58%), Gaps = 9/343 (2%) pir  C69813 RNA helicase homolog yfmL - Bacillus subtilis dbj  BAA22326.1  (D86417) YfmL [Bacillus subtilis] emb  CAB12572.1  (Z99108) similar to RNA helicase [Bacillus subtilis] Length = 376

339	

	No Hits found No Hits found	Contig117 (25807-26352 p) Contig117 (26503-26640 p)	SA-949.1 SA-950.1	SeqID 2297 SeqID 2298
Identities = 135/648 (20%), Positives = 261/648 (39%), Gaps = 90/648 (13%), emb CAB39037.2  (AL034559) hypothetical protein, PFC0960c [Plasmodium falciparum]	40	Contig117 (23809-25740 p)	SA-948.1	SeqID 2296
Identities = 265/390 (67%), Positives = 325/390 (82%), Gaps = 1/390 (0%) splP54419 METK_BACSU S-ADENOSYLMETHIONINE SYNTHETASE (METHIONINE ADENOSYLTRANSFERASE) (ADOMET SYNTHETASE) pir  D69657 methionine adenosyltransferase (EC 2.5.1.6) -Bacillus subtilis gb AAC00242.1  (AF008220) SAM synthase [Bacillus subtilis] emb CAB15033.1  (Z99119) S-adenosylmethionine synthetase [Bacillus subtilis] Length = 400	78	Contig117 (22118-23314 p)	SA-947.1	SeqID 2295
Identities = 103/315 (32%), Positives = 169/315 (52%), Gaps = 18/315 (5%) dbj BAB05404.1  (AP001512) transcriptional repressor of the biotin operon [Bacillus halodurans] Length = 323	50	Contig117 (20998-21933 m)	SA-946.1	SeqID 2294
	No Hits found	Contig117 (20847-21017 p)	SA-945.1	SeqID 2293
Identities = 408/558 (73%), Positives = 473/558 (84%), Gaps = 6/558 (1%) gb AAF98348.1 (AF280764) DNA polymerase III gamma/tau subunits [Streptococcus pyogenes]	84	Contig117 (19070-20734 p)	SA-944.1	SeqID 2292
Identities = 66/146 (45%), Positives = 95/146 (64%) pir F82706 conserved hypothetical protein XF1230 [imported] - Xylella fastidiosa (strain 9a5c) gb AAF84040.1 AE003957_1 (AE003957) conserved hypothetical protein [Xylella fastidiosa] Length = 162	54	Contig117 (18573-19070 p)	SA-943.1	SeqID 2291
Identities = 132/207 (63%), Positives = 167/207 (79%) <u>spl</u> O32033 URK_BACSU URIDINE KINASE (URIDINE  MONOPHOSPHOKINASE) pir  G69728 uridine kinase udk -  Bacillus subtilis emb CAB14675.1  (299117) uridine kinase  [Bacillus subtilis] Length = 211	62	Contig117 (17857-18486 p)	SA-942.1	SeqID 2290



Identities = 50/177 (28%), Positives = 81/177 (45%), Gaps = 31/177 (17%) pirllB72471 hypothetical protein APE2411 - Aeropyrum pernix (strain K1) dbj BAA81426.1  (AP000064) 197aa long hypothetical protein [Aeropyrum pernix] Length = 197	Identities = 71/182 (39%), Positives = 121/182 (66%), Gaps = 3/182 (1%) pir  A69859 hypothetical protein ykoE - Bacillus subtilis emb CAA05602.1  (AJ002571) YkoE [Bacillus subtilis] emb CAB13180.1  (299110) ykoE [Bacillus subtilis] Length = 199	Identities = 139/450 (30%), Positives = 224/450 (48%), Gaps = 68/450 (15%) pir  H69858 cation ABC transporter (ATP-binding protein) homolog ykoD - Bacillus subtilis emb CAA05601.1  (AJ002571) YkoD [Bacillus subtilis] emb CAB13179.1  (299110) similar to cation ABC transporter (ATP-binding protein) [Bacillus subtilis]			Identities = 167/436 (38%), Positives = 248/436 (56%), Gaps = 44/436 (10%) dbj BAB07254.1  (AP001519) unknown [Bacillus halodurans] Length = 449	Identities = 194/549 (35%), Positives = 316/549 (57%), Gaps = 36/549 (6%) gb AAB06502.1  (U50902) relaxase [Lactococcus lactis subsp. lactis] Length = 563	Identities = 31/102 (30%), Positives = 53/102 (51%), Gaps = 2/102 (1%) gb AAF72356 1 AF192329_17 (AF192329) unknown [Enterococcus faecalis] Length = 109		Identities = 54/98 (55%), Positives = 68/98 (69%), Gaps = 4/98 (4%) emb CAA69725.1  (Y08498) aggregation promoting protein [Lactobacillus gasseri]
34	63	55	No Hits found	No Hits found	98	56	40	No Hits found	98
Contig117 (26745-27314 p)	Contig117 (27311-27865 p)	Contig117 (27869-29071 p)	1. 1	Contig107 (1924-2223 m)	Contig107 (2234-3481 m)	Contig107 (3478-5109 m)	Contig107 (5081-5455 m)	Contig107 (5458-6021 m)	Contig131 (23828-24307 p)
SA-951.1	SA-952.1	SA-953.1	SA-954.1	SA-955.1	SA-956.1	SA-957.1	SA-958.1	SA-959.1	SA-96.1
SeqID 2299	SeqID 2300	SeqiD 2301	Seq1D 2302	SeqID 2303	SeqID 2304	SeqID 2305	SeqID 2306	SeqID 2307	SeqID 2308

SeqID 2309	SA-960.1	Contig107 (6024-6353 m)	No Hits found	
SeqID 2310	SA-961.1	Contig107 (6364-6663 m)	No Hits found	
SeqID_23.11	. SA-964.1	Contig107 (6715-9951 m)		Identities = 106/344 (30%), Positives = 175/344 (50%), Gaps = 21/344 (6%) ref[NP_047307.1  LtrC [Lactococcus lactis] pir  T43094 hypothetical protein - Lactococcus lactis plasmid pMRC01 gb AAC56005.1  (AE001272) LtrC [Lactococcus lactis] Length = 355
SeqID 2312	SA-966.1	Contig107 (9953-10318 m)	No Hits found	
SeqID 2313	SA-967.1	Contig107 (10360-12360 m)	42	Identities = 177/530 (33%), Positives = 296/530 (55%), Gaps = 22/530 (4%) gb AAF72343.1 AF192329_4 (AF192329) TrsK-like protein [Enterococcus faecalis]
SeqID 2314	SA-968.1	Contig107 (12405-12896 m)	32	Identities = 37/114 (32%), Positives = 56/114 (48%), Gaps = 10/114 (8%) pir E72215 oligopeptide ABC transporter, periplasmic oligopeptide-binding protein - Thermotoga maritima (strain MSB8) gb AAD36811.1 AE001813_3 (AE001813) oligopeptide ABC transporter, periplasmic oligopeptide-binding protein [Thermotoga maritima] Length = 642
SeqID 2315	SA-969.1	Contig107 (12918-13700 m)	34	Identities = 47/187 (25%), Positives = 84/187 (44%), Gaps = 21/187 (11%) pir  S45089 hypothetical protein eta - Streptococcus pyogenes plasmid pDB101 pir  S68123 hypothetical protein eta - Streptococcus pyogenes plasmid pMD101 emb CAA47095.1  (X66468) orf eta [Streptococcus pyogenes] Length = 231
SeqID 2316	SA-970.1	Contig107 (13700-13972 m)	No Hits found	
SeqID 2317	SA-971.1	Contig107 (13992-14597 m)	No Hits found	
SeqID 2318	SA-973.1	Contig107 (14618-17308 m)	2	Identities = 54/133 (40%), Positives = 66/133 (49%), Gaps = 9/133 (6%) gb AAC61959.1  (AF051917) putative membrane protein TraG [Staphylococcus aureus] prf  2004267H traG protein [Staphylococcus sp.]
SeqID 2319	SA-974.1	Contig107 (17287-17871 p)	No Hits found	



SA-975:1	—Contig107 (18065-20356 m)	20	Identities = 233/769 (30%), Positives = 387/769 (50%), Gaps = 29/769-(3%)-gb AAF72347.1 AF192329_8 (AF192329) TrsE-like protein [Enterococcus faecalis] Length = 799
SA-976.2	Contig114 (16174-17361 m)	33	Identities = 77/279 (27%), Positives = 136/279 (48%), Gaps = 14/279 (5%) splQ54955 CIAH_STRPN SENSOR PROTEIN CIAH pir  S49545 histidine kinase - Streptococcus pneumoniae emb CAA54466.1  (X77249) histidine kinase [Streptococcus pneumoniae] emb CAB54565.1  (AJ005926) histidine kinase [Streptococcus pneumoniae] Length = 444
SA-977.1	Contig114 (14493-16028 m)	06	Identities = 403/511 (78%), Positives = 466/511 (90%) gb AAC05774.1  (AF051356) D-alanine-D-alanyl carrier protein ligase [Streptococcus mutans] Length = 516
SA-978.1	Contig114 (13231-14496 m)	14	Identities = 296/418 (70%), Positives = 351/418 (83%), Gaps = 1/418 (0%) gb AAC05775.1  (AF051356) integral membrane protein [Streptococcus mutans] Length = 420
SA-98.1	Contig131 (22758-23552 p)	29	Identities = 133/263 (50%), Positives = 191/263 (72%) pir  F69742 hypothetical protein ybaF - Bacillus subtilis emb CAB11923.1  (299104) ybaF [Bacillus subtilis] Length = 265
SA-980.1	Contig114 (12977-13216 m)	. 92	Identities = 65/79 (82%), Positives = 74/79 (93%) gb AAC05776.1  (AF051356) D-alanyl carrier protein [Streptococcus mutans] gb AAC29040.1  (AF050517) unknown [Streptococcus mutans] gb AAC06286.1  (AF049357) Glg3 [Streptococcus mutans]
SA-981.1	Contig114 (11722-12984 m)	72	Identities = 238/421 (56%), Positives = 308/421 (72%), Gaps = 1/421 (0%) gb AAC29041.1  (AF050517) unknown [Streptococcus mutans]
SA-982.1	Contig114 (11155-11472 m)	No Hits found	
SA-983.1	Contig114 (9863-11155 m)	19	Identities = 49/185 (26%), Positives = 85/185 (45%), Gaps = 46/185 (24%) dbj BAB01041.1  (AB022220) gene_id:MLN21.14~unknown protein [Arabidopsis thaliana] Length = 507

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	Identities = 115/254 (45%), Positives = 172/254 (67%) sp P37545 YABD_BACSU PUTATIVE DEOXYRIBONUCLEASE YABD pir  S66068 conserved hypothetical protein yabD - Bacillus subtilis dbj BAA05274.1  (D26185) unknown [Bacillus subtilis] emb CAB11815.1  (299104) similar to hypothetical proteins [Bacillus subtilis]	Identities = 82/179 (45%), Positives = 117/179 (64%), Gaps = 4/179 (2%) splP37547 YABF_BACSU HYPOTHETICAL 20.7 KD PROTEIN IN METS-KSGA INTERGENIC REGION pir  S66070 conserved hypothetical protein yabF - Bacillus subtilis dbj BAA05276.1  (D26185) unknown [Bacillus subtilis] emb CAB11817.1  (299104) similar to hypothetical proteins [Bacillus subtilis]	Identities = 39/121 (32%), Positives = 60/121 (49%), Gaps = 12/121 (9%) pir JA83524 hypothetical protein PA0959 [imported] - Pseudomonas aeruginosa (strain PAO1) gb AAG04348.1 AE004530_1 (AE004530) hypothetical protein [Pseudomonas aeruginosa] Length = 209
No Hits found	55	09	39
Contig114 (9480-9872 m)	1	Contig114 (7672-8241 m)	Contig114 (7219-7668 m)
SA-984.1 SA-985.1	SA-986.2	SA-987.2	SA-988.2
SeqID 2329	SeqID 2331	SeqID 2332	SeqID 2333



Gaps = NOSINE NOSINE N-S RRNA STANCE Genine-Bacillus mycin 1104)	ss = ein)	.5 KD .11 tical .7.1
4%)Ga ADENOS N (16S RF (ESISTA SIN le (adeni Bac Basgamyc (299104	7%), Ga ding prot 410984.1 22.1 (Z9 ) [Bacillu	2%), Gap TCAL 31 TCAL 31 SAR11703 AA11703 I hypothe CAB159 Is subtilis
Positives = 215/284 (74% A_BACSU DIMETHYLAD DSYLMETHIONINE-6-N, THYLTRANSFERASE) (17 // EL KASUGAMYCIN RESKSGA) (KASUGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN RESKSGAMYCIN	es = 188/242 (77%), Gapporter (ATP-binding protesubtilis dbj BAA10984.1]   emb CAB11922.1  (299-binding protein) [Bacillus ngth = 276	= 147/278 (5, J HYPOTHET J HYPOTHET origilh70078 continuities dbj[B, YEBH_ECOLI subtilis] embloteins [Bacilli oteins [Bacilli r to hypothetic
tives = 2 ACSU DII TRATHIC TRANSICA (ASUGA (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (KASI (A) (A) (A) (A) (A) (A) (A) (A) (A) (A)	tives = 188/24 nsporter (ATF subtilis db lis] emb[CAB IP-binding pr Length = 276	ives = 14 CSU HY N pir[]H Ilus subti rot:YEBI al proteir milar to h
%), Positiv (SGA_BA( SGA_BA( SGA_BA( METHYL LEVEL K KSGA RASE) pir rase (EC) 7.1 (D26 3020	%), Posi ABC tra illus illus subti oorter (A	%), Posit XJB_BA 3 REGIC 3 REGIC 5 Swiss cal [Baci ypothetic 99124) si
= 155/284 (54 ) splP37468 k RASE (S-AD YL(RNA) DI YLTRANSEI imethyltransfe dbj BAA0527 nce [Bacillus sienosine trans	I/242 (58%)   E69742 A aE - Bacill wn [Bacillis] SC transpo	4/278 (33%), Popper (33%), Pop
Identities = 155/284 (54%), Positives = 215/284 (74%)Gaps.= 2/284 (0%) splP37468 KSGA_BACSU DIMETHYLADENOSINE TRANSFERASE (S-ADENOSYLMETHIONINE-6-N, N-ADENOSYL(RRNA) DIMETHYLTRANSFERASE) (16S RRNA DIMETHYLASE) (HIGH LEVEL KASUGAMYCIN RESISTANCE PROTEIN KSGA) (KASUGAMYCIN DIMETHYLTRANSFERASE) pir  S66071 probable (adenine-NG,N6-)-dimethyltransferase (EC 2.1.1) ksgA - Bacillus subtilis dbj BAA05277.1  (D26185) high level kasgamycin resistance [Bacillus subtilis] emb CAB11818.1  (299104) dimethyladenosine transferase [Bacillus subtilis] Length = 292	Identities = 141/242 (58%), Positives = 188/242 (77%), Gaps = 1/242 (0%) pir  E69742 ABC transporter (ATP-binding protein) homolog ybaE - Bacillus subtilis dbj  BAA10984.1  (D64126) unknown [Bacillus subtilis] emb  CAB11922.1  (Z99104) similar to ABC transporter (ATP-binding protein) [Bacillus subtilis]	ies = 9 5%) sp 5%) sp OTEIN hetical hor (6) hor (723) s CAB16
2/284 - 1dentit 2/284 - 1Z284	1/242 1/242 hor (D6412) sim	Identiti 16/278 ( PR hypot (D8302 protein; (Z99 embli
72	63	. 51
. 22	<b>o</b>	S.
2 m)	(65 p)	(m.g.
(6320-7192 m)	(21923-22765 p)	Contig114 (5465-6316 m)
Contig114 (6	Contig131 (21	tig114 (5
CO	Cont	Cou
SA-989.1	SA-99.1	SA-990.1
SeqID 2334	SeqID 2335	SeqID 2336
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SeqID 2337	SA-991.1	Contig114 (4437-5309 m)	69	Identities = 149/296 (50%). Positives = 204/296 (68%), Gaps = 14/296 (4%) pir. A65879 conserved hypothetical protein yloQ - Bacillus subtilis emb[CAA74251.1  (Y13937) YloQ protein [Bacillus subtilis] emb[CAB13451.1  (Z99112) similar to hypothetical proteins [Bacillus subtilis]
SeqID 2338	SA-993.1	Contig114 (3768-4430 m)	. 89	Identities # 112/211 (53%), Positives = 152/211 (71%) dbj BAB06221.1  (AP001515) unknown conserved protein [Bacillus halodurans] Length = 216
SeqID 2339	SA-994.1	Contig114 (3143-3775 m)	52	Identities = 75/220 (34%), Positives = 112/220 (50%), Gaps = 18/220 (8%) pirl C69879 hypothetical protein yloS - Bacillus subtilis emb CAA74253.1  (Y13937) YloS protein [Bacillus subtilis] emb CAB13453.1  (299112) yloS [Bacillus subtilis] Length = 214
SeqID 2340	SA-995.1	Contig114 (1867-3099 m)	54	Identities = 157/393 (39%), Positives = 238/393 (59%), Gaps = 3/393 (0%) pirl D82810 conserved hypothetical protein XF0413 [imported] - Xylella fastidiosa (strain 9a5c) gb AAF83223.1 AE003892_8 (AE003892) conserved hypothetical protein [Xylella fastidiosa] Length = 456
SeqID 2341	SA-996.1	Contig114 (936-1877 m)	. 19	Identities = 139/313 (44%), Positives = 196/313 (62%), Gaps = 6/313 (1%) gblAAC44803.1  (U21636) cmp-binding-factor 1 [Staphylococcus aureus]
SeqID 2342	SA-997.1	Contig114 (27-839 m)	7.1	Identities = 142/270 (52%), Positives = 196/270 (72%), Gaps = 1/270 (0%) emb[CA410902.1] (AJ222642) purR [Lactococcus lactis] Length = 271
SeqID 2343	SA-998.2	Contig127 (15315-15944 m)	30	Identities = 47/120 (39%), Positives = 69/120 (57%), Gaps = 5/120 (4%) dbj BAA11325.1  (D78257) ORF8 [Enterococcus faecalis] Length = 120

-				Identities = 98/265 (36%), Positives = 161/265 (59%), Gaps = 4/265 (1%) splO66553 PROC_AQUAE PYRROLINE-5-
SeqID 2344	SA-999.1	Contig127 (14524-15294 p)	69	CARBOXYLATE REDUCTASE (P5CR) (P5C REDUCTASE) pirl F70315 pvrroline carboxylate reductase - Aquifex aeolicus
!			į	gbjAAC06504.1  (AE000676) pyrroline carboxylate reductase [Aquifex aeolicus] Length = 265



TABLEAU 2. Protéines de surface de Streptococcus agalactiae

	SEQ ID	N° IPF	Motif LPXTG	% identité/ similitude
1	886	18	LPHTG	24/37 : Sec (Surface exclusion protéine) E. faecalis
2	386	1250	LPSTG	22/40 :Sec10 (Surface exclusion protéine)
		1230	L	Enterococcus faecalis plasmid pCF10
3	1669	280	LPKTG	50/65: pullulanase S. pneumoniae
4	1895	523	LPATG	90/91 : alpha like protéine S. agalactiae
5	806	1716	LPSTG	31/46: M-like protéine S. equi
6	1183	2140	LPLTG	36/60: cell-division protein homolog ywcF B. subtilis 34/56: stage V sporulation protein E B. halodurans
7	1346	2337	LPKTG	25/38 : Unknown B. halodurans
8	1942	571	LPSTG	50/60: Hemagglutinine S. gordonii
9	2129	765	LPNTG	28/43 : sdrc <i>S. aureus</i>
10	2046	678	LPKTG	57/70: putative cyclo-nucleotide phosphodiesterase  Strep. dysgalactiae susp. equismilis
11	618	1503	LPKTG	49/65 : celle envelope proteinase S. thermophilus
	1227	220	LPSTG	30/43 : SspB (adhesion) S. gordonii
	1954	584	LPKTG	26/48: hypothetical serine riche repeat prot S. pombe
	1493	2495	LPKTG	23/47 : gene drosophile
				30/46 : 6-aminohexanoate-cyclic-dimer hydrolase
15	1955	585	LPKTG	Deinococcus radiodurans.
16	943	1861	LPKTG	50/62: hypothetical protein 2 S. mutans
	1221	2192	LPKTG	35/48 : C. elegans UNC-89 (6642 aa)
18	613	15	LPSTG	37/52: SpaA: Ag de surface de S. sobrinus
19	382	1247	LPSTG	38/52: SpaA: Ag de surface de S. sobrinus
20	792	17	LPKTG	30 /47 : hypothetical protéine iota de S. pyogènes plasmid pDB101
21	383	1248	LPRTG	38/52 : hypothetical protéine iota de S. pyogènes plasmid pDB101
22	1418	2414	LPSTG	33/47 : orfC E. faecalis, plasmid pAM-beta -1
	·			protéines possibles (IPXTG, FPXTG)
				31/43 : Inconnue B. Halodurans
23	2171	807	IPQTG	29/41 : fimbrial SU Actinomyces naeslundii
23	21/1	<b>0</b> 0 /	irQiG	24:41: putative cell-surface adhesin SdrF
L				[Staphylococcus epidermidis]
				27/38 : hypothetical protein 2 <i>L. leichmanii</i>
24	1343	2334	IPQTG	23/38 : fimbrial SU Actinomyces naeslundii
				, , , , , , , , , , , , , , , , , , ,
25	659	1551	FPKTG	37/50: fibronectin binding protein I S. pyogenes



TABLE	AU 3: Liste d	es phases co	dantes annotées	identifiées par l'analyse d	TABLEAU 3: Liste des phases codantes annotées identifiées par l'analyse de la séquence génomique complète SEQ ID N°2345
		Anciens			
	SEQ ID	SEQ ID		Positions / Sens	
SEQ ID (ADN)	(Protéine)	(Protéine)	Protéine N°	(p=plus, m=minus)	Fonctions Prédites
SEO ID 2º 4482	2766	7.4.0	2 4000 4	306600 000000	similar to glutamyl-aminopeptidase and/or to endo-1,4-beta-
SEG ID 11 4402	2340	147	3A-1000.1	T02238-203305 m	giucanase
SEQ ID n° 4483	2347	143	SA-1001.1	203490-203729 m	Unknown
SEQ ID n° 4484	2348	144	SA-1002.1	203890-204174 p	similar to unknown protein
SEQ ID n° 4485	2349	145	SA-1003.1	204171-204494 p	similar to thioredoxin H1
SEQ ID n° 4486	2350	146	SA-1004.1	204527-205153 p	weakly similar to phenylalanyl-tRNA synthetase (beta subunit)
SEQ ID n° 4487	2351	147	SA-1006.1	205207-205923 m	similar to unknown proteins
SEQ ID n° 4488	2352	148	SA-1007.1	206004-206399 p	similar to single-strand binding protein
SEQ ID n° 4489	2353	149	SA-101.1	2190539-2191378 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4490	2354	150	SA-1010.1	206522-207166 p	similar to unknown proteins
SEQ 1D n° 4491	2355	151	SA-1012.1	207193-208938 p	Similar to two-component sensor histidine kinase
SEQ ID n° 4492	2356	152	SA-1013.1	208919-209659 p	Similar to two-component response regulator
SEQ ID n° 4493	2357	154	SA-1016.1	209829-210284 p	similar to unknown proteins
SEQ ID n° 4494	2358	155	SA-1017.1	210286-211014 p	similar to unknown proteins
SEQ ID n° 4495	2359	156	SA-1018.1	211257-212885 p	Similar to (oligopeptide) ABC transporter (binding protein)
SEQ ID n° 4496	2360	157	SA-1019.1	212998-213975 p	Similar to oligopeptide ABC transporter (permease)
SEQ ID n° 4497	2361	158	SA-102.1	2191378-2191917 m	similar to phosphatidylglycerophosphate synthase
SEQ ID n° 4498	2362	159	SA-1020.1	213972-214793 p	Similar to ABC transporter (permease)
SEQ ID n° 4499	2363	160	SA-1021.1	214805-215608 p	Similar to oligopeptide ABC transporter (ATP-binding protein)
SEQ ID n° 4500	2364	161	SA-1022.2	215592-216218 p	Similar to oligopeptide ABC transporter (ATP-binding protein)
SEQ ID n° 4501	2365	163	SA-1024.2	10743-10877 p	Hypothetical protein :
SEQ ID n° 4502	2366	164	SA-1025.2	10369-10740 p	similar to unknown protein
SEQ ID n° 4503	2367	165	SA-1026.1	10110-10382 p	similar to unknown protein
SEQ ID n° 4504	2368	166	SA-1028.1	6322-9819 p	similar to transcription-repair coupling factor
SEQ ID n° 4505	2369	167	SA-1029.1	5750-6325 p	similar to peptidyl-tRNA hydrolase
SEQ ID n° 4506	2370	168	SA-103.1	2192043-2193326 m	similar to unknown proteins
SEQ ID n° 4507	2371	169	SA-1030.1	4551-5666 p	similar to hypothetical GTP binding protein
SEQ ID n° 4508	2372	170	SA-1031.1	4188-4391 p	Hypothetical protein
SEQ ID n° 4509	2373	171	SA-1032.1	3789-3986 p	similar to unknown protein
SEQ ID n° 4510	2374	172	SA-1033.1	2898-3779 p	similar to putative transcription regulator
SEQ ID n° 4511	2375	173	SA-1034.1	1692-2828 p	beta subunit of DNA polymerase III

SEQ ID nº 4512	2376	174	SA-1035.2	176-1537 p	replication initiation protein DnaA
SEQ ID n° 4513	2377	175	SA-1036.2	2217128-2217901 p	similar to chromosome partitioning protein ParB
SEQ ID n° 4514	2378	176	SA-1037.1	2215801-2217030 p	similar to serine protease
SEQ ID nº 4515	2379	177	SA-1038.1	2215121-2215600 m	similar to unknown protein
SEQ ID n° 4516	2380	178	SA-1039.1	2211963-2214542 p	similar to unknown transmembrane protein
SEQ ID n° 4517	2381	179	SA-104.1	2193328-2194572 m	similar to unknown proteins
SEQ ID n° 4518		180	SA-1041.1	2210221-2211840 p	similar to ABC transporter (ATP-binding protein) ····
SEQ ID n° 4519	2383	181	SA-1042.4	2209283-2210155 p	similar to unknown proteins
SEQ ID n° 4520	2384	182	SA-1044.2	891360-892031 p	similar to thiamine-phosphate pyrophosphorylase (ThiE)
SEQ ID n° 4521	2385	183	SA-1045 1	892158-893417 p	Similar to UDP-N-acetylolucosamine 1-carboxovinyltransferase
SEQ ID n° 4522	2386	184	SA-1046.1	893519-894073 p	Similar to acetyl transferase
SEQ ID n° 4523	2387	185	SA-1047.1	894066-895349 p	Similar to unknown proteins
SEQ ID n° 4524	2388	186	SA-1048.1	895372-896232 p	similar to methionine aminopeptidase
SEQ ID n° 4525	2389	187	SA-1049.1	896234-897154 p	similar to unknown proteins
SEQ ID n° 4526.	2390	188	SA-105.1	2194811-2195164 p	similar to unknown proteins
SEQ ID n° 4527	2391	189	SA-1050.1	897171-897626 m	similar to unknown proteins
SEQ ID n° 4528	2392	190	SA-1051.1	897808-898317 p	similar to unknown proteins
SEQ ID n° 4529	2393	191	SA-1053.1	898457-900415 p	similar to DNA ligase
SEQ ID n° 4530	2394	192	SA-1054.1	900427-901446 p	similar to unknown proteins
SEQ ID n° 4531	2395	193	SA-1055.1	901450-903750 p	Similar to pullulanase
SEQ ID n° 4532	2396	194	SA-1056.1	903956-905824 p	similar to 1,4-alpha-glucan branching enzyme
SEQ ID n° 4533	2397	195	SA-1057.1	905866-907005 p	similar to glucose-1-phosphate adenylyltransferase
SEQ ID n° 4534	2398	196	SA-1058.1	906995-908128 p	similar to glucose-1-phosphate adenylyltransferase
SEQ ID n° 4535	2399	197	SA-1059.2	908125-909555 p	similar to glycogen synthase
SEQ ID n° 4536	2400	198	SA-106.1	2195167-2196276 p	Similar to recombination protein RecF
SEQ ID n° 4537	2401	201	SA-1062.2	1167875-1169149 m	similar to xanthine permease
SEQ ID n° 4538	2402	202	SA-1063.1	1166988-1167776 p	similar to nitrite or formate transporter
SEQ ID n° 4539	2403	203	SA-1064.1	1165654-1166889 p	similar to unknown protein (N-terminal part)
SEQ ID n° 4540	2404	204	SA-1065.1	1165028-1165630 p	Uniknown, similar to unknown protein
					similar to other protein (putative thiamine biosynthesis
SEQ ID n° 4541	2405	205	SA-1066.1	1164072-1165010 p	(lipoprotein)
SEQ ID n° 4542	2406	206	SA-1067.1	1163741-1163923 p	similar to 4-oxalocrotonate tautomerase
SEQ ID n° 4543	2407	207	SA-1068.1	1163034-1163603 m	thymidine kinase
SEQ ID n° 4544	2408	208	SA-1069.1	1161920-1162999 m	peptide chain release factor 1
SEQ ID n° 4545	2409	500	SA-107.1	2196288-2197154 m	putative transmembrane protein similar to unknown protein



SEQ ID n° 4546	2410	210	SA-1071.1	1161090-1161920 m	similar to protoporphyrinogen oxidase
SEQ ID n° 4547	2411	211	SA-1072.1	1160501-1161097 m	similar to unknown protein
₽	2412	212	SA-1073.1	1159153-1160409 m	serine hydroxymethyltransferase
Ω	2413	213	SA-1074.1	1158171-1159148 m	similar to unknown protein
ଠା	2414	214	SA-1075.1	1157567-1158169 m	similar to unknown protein
ଠା	2415	215	SA-1076.1	1155834-1157555 m	similar to ABC transporter (ATP-binding protein)
₽	2416	216-	SA-1077.1	1154100-1155833 m	similar to-ABC transporter (ATP-binding protein)
의	2417	217	SA-1078.2	1153889-1154086 m	Unknown
₽	2418	218	SA-1079.1	1152107-1153825 p	similar to phosphoglucomutase
	2419	219	SA-108.1	2197210-2197872 m	similar to transcriptional regulator
SEQ ID n° 4556	2420	220	SA-1080.1	1151425-1151997'p	similar to unknown protein
SEQ ID n° 4557	2421	221	SA-1081.1	1150837-1151379 p	similar to pantothenate metabolism flavoprotein
SEQ ID n° 4558	2422	222	SA-1082.1	1150158-1150844 p	similar to flavoprotein
SEQ ID n° 4559	2423	223	SA-1083.1	1149168-1150004 m	similar to unknown protein
SEQ ID n° 4560	2424	224	SA-1084.1	1147976-1149175 m	similar to NADH-dependent oxidoreductase
SEQ ID n° 4561	2425	225	SA-1085.1	1146993-1147979 m	similar to unknown protein
SEQ ID n° 4562	2426	226	SA-1086.1	1146632-1146964 m	similar to glycine cleavage system H protein
SEQ ID n° 4563	2427	227	SA-1087.2	1145821-1146639 m	similar to unknown protein
SEQ ID n° 4564	2428	228	SA-1088.2	346400-347701 p	Similar to cellobiose-specific PTS enzyme IIC
SEQ ID n° 4565	2429	229	SA-1089.1	346078-346398 p	Similar to cellobiose-specific PTS enzyme IIB
SEQ ID n° 4566	2430	230	SA-109.1	2197947-2199428 m	similar to inosine monophosphate dehydrogenase
SEQ ID n° 4567	2431	231	SA-1090.1	345738-346061 p	Similar to cellobiose-specific PTS enzyme IIA
SEQ ID n° 4568	2432	232	SA-1091.2	344551-345534 p	similar to hypothetical transcriptional regulator
SEQ ID n° 4569	2433	233	SA-1092.2	343677-344432 p	similar to transcriptional regulator (DeoR family)
SEQ ID n° 4570	2434	234	SA-1093.1	342780-343556 m	similar to pyruvate formate-lyase activating enzyme
SEQ ID n° 4571	2435	235	SA-1094.1	342377-342733 p	similar to general stress protein GSP13
SEQ ID n° 4572	2436	236	SA-1095.1	340975-342375 p	C-terminal part similar to peptidyl-prolyl cis-trans isomerase
SEQ ID n° 4573	2437	237	SA-1096.1	340292-340933 p	Similar to two-component response regulator
SEQ ID n° 4574	2438	238	SA-1097.1	339280-340299 p	Similar to two-component sensor histidine kinase
SEQ ID n° 4575	2439	239	SA-1098.1	, 338588-339283 p	similar to unknown proteins
SEQ.ID n° 4576	2440	241	SA-11.1	1031773-1034922 m	similar to plasmid proteins
SEQ ID n° 4577	2441	242	SA-110.1	2199585-2200070 m	similar to arginine repressor (ArgR)
SEQ ID n° 4578	2442	243	SA-1100.1	336474-338429 p	similar to hypothetical protein kinases
SEQ ID n° 4579	2443	244	SA-1101.1	335737-336474 p	similar to other proteins
SEQ ID n° 4580	2444	245	SA-1102.1	334377-335699 p	similar to RNA-binding protein Sun
SEQ ID n° 4581	2445	246	SA-1103.1	333452-334387 p	methionyl tRNA formyltransferase

SEQ ID n° 4582 SEQ ID n° 4583	2446	247	SA-1104.1 SA-1105.2	331015-333405 p	primosomal replication factor Y (primosomal protein N ) similar to unknown proteins
SEQ ID nº 4584	2448	249	SA-1107.2	141808-143196 p	similar to argininosuccinate Ivase
SEQ ID n° 4585	2449	250	SA-1108.1	140599-141789 p	similar to argininosuccinate synthase
SEQ ID n° 4586	2450	251	SA-1109.1	139375-140445 p	similar to two-component sensor histidine kinase
SEQ ID n° 4587	2451	252	SA-111.1	2200080-2200760 m	similar to transcriptional regulator (Crp/Fnr family)
SEQ ID n° 4588	2452	253	SA-1110.1	. 138711-139382 p	similar to similar to two-component response regulator
SEQ ID n° 4589	2453	254	SA-1111.1	137994-138680 p	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4590	2454	255	SA-1112.1	136906-137994 p	similar to putative ABC transporter (permease)
SEQ ID n° 4591	2455	256	SA-1113.1	135738-136724 m	similar to transcription repressor of ribose operon
SEQ ID n° 4592	2456	257	SA-1114.1	134834-135745 m	similar to ribokinase
SEQ ID n° 4593	2457	258	SA-1115.1	134461-134859 m	similar to ribose ABC transporter (permease)
SEQ ID n° 4594	2458	259	SA-1116.1	132967-134445 m	similar to ribose ABC transporter (ATP-binding protein)
SEQ ID n° 4595	2459	260	SA-1118.1	132024-132965 m	similar to ribose ABC transporter (permease)
SEQ ID n° 4596	2460	261	SA-1119.1	131003-131971 m	similar to ribose ABC transporter (binding protein)
SEQ ID n° 4597	2461	262	SA-112.1	2200968-2201672 p	similar to unknown proteins
SEQ ID n° 4598	2462	263	SA-1120.1	129509-130963 p	glutamyl-tRNA synthetase
SEQ ID n° 4599	2463	264	SA-1121.1	127981-129300 p	similar to mercury(II) reductase
SEQ ID n° 4600	2464	265	SA-1122.1	127330-127827 p	similar to carbonic anhydrase
SEQ 1D n° 4601	2465	266	SA-1123.2	125830-127194 p	similar to DNA repair protein RadA
					similar to N-acetylgalactosamine-specific PTS system enzyme
SEQ ID n° 4602	2466	268	SA-1125.2	1963678-1964112 m	IIA
SEQ ID nº 4603	2467	269	SA-1126.2	1962479-1963675 m	similar to glucuronyl hydrolase
					similar to N-acetylgalactosamine-specific PTS system enzyme
SEQ 1D n° 4604	2468	270	SA-1127.1	1961930-1962424 m	·IIA
					similar to N-acetylgalactosamine-specific PTS system enzyme
SEQ ID n° 4605	2469	271	SA-1129.1	1961028-1961894 m	110
SEQ ID n° 4606	2470	272	SA-113.1	2201939-2203171 p	similar to arginine deiminase
					similar to N-acetylgalactosamine-specific PTS system enzyme
SEQ ID n° 4607	2471	273	SA-1131.1	1960226-1961041 m	OII
0000	273	71.0	· · · · · · · · · · · · · · · · · · ·	40E0242 40E044E ==	initiate to henciting autote lease from Flauchacterium hencitaires
3EQ ID II 4000	7/47	4/7	3A-1132.1	1996242-1900140111	פוווווומן כל ווכלימוויוון פתומים ולמפס ווכווון ומיכסמכים ומווון ומיכסמכים ומווון ומיכסמכים ומינון ומיכסמכים ומיכון ומיכסמכים ומיכון ומיכסמכים ומיכון ומיכון ומיכסמכים ומיכון ומי
SEQ ID n° 4609	2473	275	SA-1134.1	1957161-1958162 m	similar to transcriptional regulator (Laci family)
SEQ ID n° 4610	2474	276	SA-1135.1	1956476-1957090 p	similar to polypeptide deformylase
SEQ ID n° 4611	2475	277	SA-1136.1	1955766-1956410 p	similar to transcriptional regulator (Crp/Fnr family)
SEQ ID n° 4612	2476	278	SA-1137.1	1954491-1955654 m	similar to permease (major facilitator superfamily)



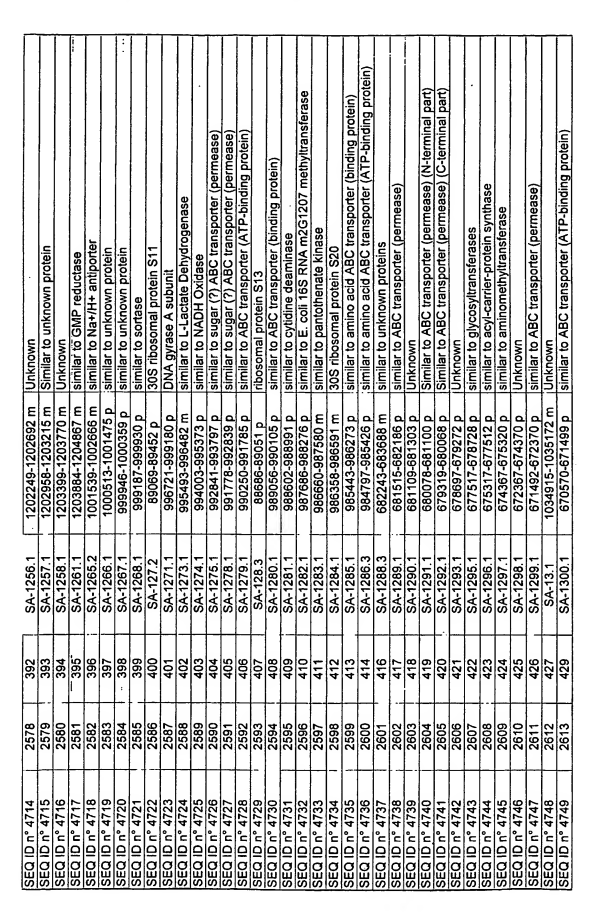
SEQ ID n° 4613	2477	279	SA-1138.1	1953462-1954445 n	Similar to oxidoreductase
SEQ ID n° 4614	2478	280	SA-1139.1	1951419-1953314 m	similar to endopeptidase
SEQ ID n° 4615	2479	281	SA-114.1	2203267-2203608 p	similar to unknown proteins
SEQ ID n° 4616	2480	282	SA-1140.1	1950416-1951369 p	similar to L, D-carboxypeptidase LdcA
SEQ ID n° 4617	2481	283	SA-1141.1	1950013-1950249 m	lysin rich protein
SEQ ID n° 4618	2482	284	SA-1143.2	1947942-1950011 m	similar to Na+/H+ antiporter
					similar to alkyl hydroperoxide reductase (large subunit) and
SEQ ID n° 4619	2483	285	SA-1144.1	1946133-1947665 p	NADH dehydrogenase
SEQ ID n° 4620	2484	286	SA-1145.2	1945555-1946115 p	similar to alkyl hydroperoxide reductase (small subunit)
SEQ ID n° 4621	2485	288	SA-1148.2	1764745-1765479 m	similar to unknown proteins
SEQ ID n° 4622	2486	289	SA-1149.1	1764471-1764722 m	Unknown
SEQ ID n° 4623	2487	290	SA-115.1	2203624-2204637 p	similar to ornithine transcarbamylase
SEQ ID n° 4624	2488	291	SA-1150.1	1763365-1764474 m	Similar to unknown proteins
SEQ ID n° 4625	2489	292	SA-1151.1	1762559-1763272 m	Similar to transcriptional regulator
SEQ ID n° 4626	2490	293	SA-1152.1	1762016-1762420 m	Similar to unknown proteins
SEQ ID n° 4627	2491	294	SA-1153.1	1761037-1761750 p	Similar to glycerol uptake facilitator
SEQ ID n° 4628	2492	295	SA-1154.1	1760653-1761027 p	Similar to unknown proteins
SEQ ID n° 4629	2493	296	SA-1155.1	1760075-1760653 p	Similar to unknown proteins
SEQ ID n° 4630	2494	297	SA-1156.1	1759048-1760037 p	Similar to hypothetical dihydroxyacetone kinase
SEQ ID n° 4631	2495	298	SA-1157.1	1758338-1758874 m	Similar to putative transcription regulator
SEQ ID n° 4632	2496	299	SA-1158.1	1757342-1758328 m	Similar to putative dihydroxyacetone kinase
SEQ ID n° 4633	2497	300	SA-116.1	2204700-2206127 p	similar to arginine/ornithine antiporter
SEQ ID n° 4634	2498	301	SA-1160.1	1756508-1757224 m	Similar to unknown proteins
SEQ ID n° 4635	2499	302	SA-1161.1	1755664-1756353 m	Similar to other proteins
SEQ ID n° 4636	2500	303	SA-1162.1	1754729-1755562 m	similar to (amino acid??) ABC transporter (binding protein)
SEQ ID n° 4637	2501	304	SA-1163.1	1753764-1754588 m	Similar to unknown protein
SEQ ID n° 4638	2502	305	SA-1165.1	1752254-1753630 m	Similar to unknown proteins
SEQ ID n° 4639	2503	307	SA-1167.1	1751191-1752261 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4640	2504	308	SA-1169.1	1750502-1751194 m	similar to ABC transporter (permease)
SEQ ID n° 4641	2505	309	SA-117.1	2206148-2207104 p	similar to carbamate kinase
SEQ ID n° 4642	2506	310	SA-1170.1	1749398-1750453 p	similar to NAD alcohol dehydrogenase
SEQ ID n° 4643	2507	311	SA-1171.2	1747917-1749284 p	Similar to branched-chain amino acid transporter
SEQ ID n° 4644	2508	312	SA-1173.3	1370419-1370727 m	truncated C-terminal part
SEQ ID n° 4645	2509	313	SA-1174.1	1370724-1371032 m	Unknown
SEQ ID n° 4646	2510	314	SA-1175.1	1371034-1371606 m	fruncated C-terminal part
SEQ ID n° 4647	2511	315	SA-1176.1	1371607-1372110 m	truncated N-terminal part

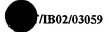
			zyme					IIBC	) IIA				arge subunit)	nnit)						protein												tein		(No.
Similar to unknown proteins	Unknown	tryptophanyl-tRNA synthetase	similar to probable type II DNA modification enzyme	similar to integrase	similar to unkhown proteins	similar to unknown proteins	similar to Phospho-beta-D-galactosidase	Similar to lactose-specific PTS system enzyme IIBC	Similar to lactose-specific PTS system enzyme IIA	Similar to transcription antiterminator	similar to tagatose-1,6-diphosphate aldolase	similar to tagatose 6-phosphate kinase	similar to galactose-6-phosphate isomerase (large subunit)	similar to galactose 6-P isomerase (small subunit)	Unknown	similar to unknown proteins	similar to unknown proteins	similar to unknown protein	Unknown	similar to plasmid recombination / mobilization protein	Unknown	similar to plasmid replication protein	Unknown	similar to replication protein (N-terminal part)	Unknown	Unknown	similar to transcriptional regulator	similar to integrase	ribosomal protein S9	50S ribosomal protein L13	similar to unknown proteins	similar to N-acetylglucosamine metabolism protein	similar to other proteins	(Mangarana III appointment and a reliais
1372659-1373864 m					-		1381531-1382937 m		1384730-1385047 m			1387287-1388216 m	1388229-1388744 m	1388761-1389186 m	248617-249024 p	_	247430-247717 m	247105-247440 m											238812-239204 p				949709-950446 p	
SA-1178.1	SA-1179.1	SA-118.2	SA-1180.2	SA-1182.1	SA-1183.1	SA-1184.1	SA-1187.1	SA-1189.1	SA-1190.1	SA-1191.1	SA-1192.1	SA-1193.1	SA-1194.3	SA-1195.3	SA-1196.2	SA-1197.1	SA-1198.1	SA-1199.1	SA-1201.1	SA-1203.1	SA-1205.1	SA-1207.1	SA-1208.1	SA-1209.1	SA-1210.1	SA-1211.1	SA-1212.1	SA-1213.1	SA-1214.2	SA-1215.2	SA-1216.2	SA-1217.1	SA-1218.1	10101
317	318	319	320	321	322	323	324	326	328	329	330	331	332	333	334	335	336	337	340	342	343	344	345	346	348	349	350	351	352	353	354	355	356	257
2513	2514	2515	2516	2517	2518	2519	2520	2521	2522	2523	2524	2525	2526	2527	2528	2529	2530	2531	2532	2533	2534	2535	2536	2537	2538	2539	2540	2541	2542	2543	2544	2545	2546	25.47
SEQ ID n° 4649	SEQ ID n° 4650	SEQ ID n° 4651	ů	n°	'n	°	°	SEQ ID n° 4657	SEQ ID n° 4658	SEQ ID n° 4659	SEQ ID n° 4660	SEQ ID n° 4661	SEQ ID n° 4662	SEQ ID n° 4663	SEQ ID n° 4664	SEQ ID n° 4665	SEQ ID n° 4666	SEQ ID n° 4667	SEQ ID n° 4668	SEQ ID n° 4669	SEQ ID n° 4670	SEQ ID n° 4671	SEQ ID n° 4672	SEQ ID n° 4673	SEQ ID n° 4674	SEQ ID n° 4675	SEQ ID n° 4676	SEQ ID n° 4677	SEQ ID n° 4678	SEQ ID n° 4679	₽	ID n°	SEQ ID n° 4682	CEO ID 20 4683



SEO ID nº 4684	2548	358	SA-1220 1	048403-048483 5	eimilar to inknown proteine
SEQ ID nº 4685	2549	359	SA-1221.1	947707-948078 p	similar to unknown proteins
SEQ ID n° 4686	2550	360	SA-1222.1	946232-947584 p	similar to phospho-sugar mutase
SEQ 1D n° 4687	2551	361	SA-1223.1	945219-946178 p	similar to unknown proteins
SEQ ID n° 4688	2552	362	SA-1225.1	944365-945216 p	similar to unknown proteins
4					similar to unknown protein (to hypothetical UDP-N-
SEQ ID n° 4689	2553	363	SA-1226.1	942882=944225 m	acetylmuramyl tripeptide synthetase???)
SEQ ID n° 4690	. 2554	364	SA-1227.1	942097-942882 m	similar to other proteins
SEQ ID n° 4691	2555	365	SA-1228.1	941001-941990 p	similar to lipoate-protein ligase A
SEQ ID n° 4692	2556	366	SA-123.1	90455-90841 p	50S ribosomal protein L17
SEQ ID n° 4693	2557	367	SA-1230.1	939146-940903 p	similar to acetoin dehydrogenase E3 component (dihydrolipoamide dehydrogenase)
SEQ 10 n° 4694	2558	368	SA-1231.1	937698-939086 p	similar to acetoin dehydrogenase E2 component (dihydrolipoamide acetyltransferase)
SEQ 1D n° 4695	2559	369	SA-1232.1	936573-937571 p	similar to acetoin dehydrogenase (TPP-dependent) beta chain
SEQ ID n° 4696	2560	371	SA-1234.1	935530-936498 p	similar to acetoin dehydrogenase (TPP-dependent) alpha chain
SEQ 1D n° 4697	2561	372	SA-1236.3	933471-935381 p	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4698	2562	373	SA-1238.2	1190234-1190593 m	Unknown
SEQ ID n° 4699	2563.	374	SA-1239.1	1190655-1191113 m	Unknown
SEQ ID n° 4700	2564	375	SA-1240.1	1191174-1191401 m	Unknown
SEQ ID n° 4701	2565	378	SA-1243.1	1191645-1192142 m	Unknown
SEQ ID n° 4702	2566	379	SA-1244.1	1192156-1192908 m	Unknown
SEQ ID n° 4703	2567	. 380	SA-1245.1	1192957-1193442 m	Unknown
SEQ ID n° 4704	2568	381	SA-1246.1	1193548-1193775 m	Unknown
SEQ ID n° 4705	2569	382	SA-1247.1	1194050-1196848 m	putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 4706	2570	383	SA-1248.1	1196915-1197625 m	putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 4707	2571	385	SA-125.1	89502-90440 p	RNA polymerase (alpha subunit)
					similar to surface exclusion protein Sec10 precursor -
SEQ ID n° 4708	2572	386	SA-1250.1	1197640-1199871 m	Enterococcus faecalis plasmid pCF10
SEQ ID n° 4709	2573	387	SA-1251.1	1199888-1200187 m	Unknown
SEQ ID n° 4710	2574	388	SA-1252.1	1200570-1200767 m	Similar to phage protein
SEQ ID n° 4711	2575	389	SA-1253.1	1200764-1200949 m	Unknown
SEQ ID n° 4712	2576	390	SA-1254.1	1200951-1201976 m	similar to putative plasmid replication initiator protein A
SEQ ID n° 4713	2577	391	SA-1255.1	1201978-1202142 m	Unknown







SEQ ID n° 4750	2614	430	SA-1301.1	670104-670580 p	similar to (3R)-hydroxymyristoyl-[acyl carrier protein] dehydratase
SEQ ID n° 4751	2615	431	SA-1302.1	669815-670120 p	similar to acyl carrier protein
SEQ ID n° 4752	2616	432	SA-1303.1	669100-669822 p	similar to 3-oxoacyl-(acyl-carrier protein) reductase
SEQ ID n° 4753	2617	433	SA-1304.2	   668255-669103 p	weakly similar to malonyl-CoA; Acyl carrier protein transacylase
SEQ ID n° 4754	2618	434	SA-1305.2	667950-668255 p	Unknown
SEQ ID n° 4755	2619	435	SA-1308.2	1846687-1847346 p	Similar to transcriptional regulator (FNR/CRP family)
SEQ ID n° 4756	2620	436	SA-1309.1	1847370-1849655 p	Similar to X-prolyl dipeptidyl aminopeptidase
SEQ ID n° 4757	2621	437	SA-1310.1	1849659-1850018 p	Similar to unknown proteins
SEQ ID n° 4758	2622	438	SA-1311.1	1850064-1851044 p	Similar to heptaprenyl diphosphate synthase component II
					Similar to cytochrome D ABC transporter ATP binding and
SEQ ID n° 4759	2623	439	SA-1312.1	1851129-1852877 m	permease protein
					Similar to cytochrome D ABC transporter (ATP binding and
SEQ 1D n° 4760	2624	440	SA-1313.1	1852870-1854588 m	permease protein)
SEQ ID n° 4761	2625	441	SA-1314.1	1854588-1855607 m	Similar to cytochrome bd ubiquinol oxidase (subunit II)
SEQ ID n° 4762	2626	442	SA-1316.1	1855608-1857035 m	Similar to cytochrome bd ubiquinol oxidase (subunit I)
SEQ ID n° 4763	2627	443	SA-1318.1	1857138-1858346 m	Similar to hypothetical NADH dehydrogenase
i .		į			
SEQ ID n° 4/64	2628	444	SA-1319.2	1858359-1859258 m	Similar to hypothetical menaquinone biosynthesis protein MenA
SEQ ID n° 4765	2629	446	SA-1320.1	1108449-1108991 m	Unknown
SEQ ID n° 4766	2630	447	SA-1322.1	1108996-1109481 m	Unknown
SEQ ID n° 4767	2631	449	SA-1324.1	1109840-1110130 m	similar to immunogenic secreted protein precursor (C-terminal part)
					•
SEQ ID n° 4768	2632	450	SA-1325.1	1110093-1110989 m	similar to immunogenic secreted protein precursor (internal part)
SEQ ID n° 4769	2633	451	SA-1326.1	1110908-1111336 m	Unknown
SEQ ID n° 4770	2634	452	SA-1327.1	1111463-1111717 m	similar to unknown protein
SEQ ID n° 4771	2635	453	SA-1328.1	1111738-1112328 m	similar to other protein
SEQ ID n° 4772	2636	454	SA-1329.1	1112342-1112647 m	Unknown
SEQ ID n° 4773	2637	456	SA-1330.1	1112776-1113690 m	similar to unknown protein
SEQ ID n° 4774	2638	457	SA-1331.1	1113693-1114049 m	Unknown
SEQ ID n° 4775	2639	458	SA-1332.1	1114061-1114411 m	similar to unknown protein
SEQ ID n° 4776	2640	460	SA-1334.1	1114425-1118396 m	weakly similar to DNA-translocase
SEQ ID n° 4777	. 2641	461	SA-1335.1	1118329-1118832 m	Unknown
SEQ ID n° 4778	2642	462	SA-1336.1	1118839-1120113 m	similar to unknown protein



SEQ ID n° 4779	2643	463	SA-1337.1	1120113-1120355 m	similar to unknown protein
SEQ ID n° 4780	2644	464	SA-1338.1	1120339-1120812 m	Unknown
SEQ ID n° 4781	2645	465	SA-1339.3	1120821-1123832 m	similar to unknown protein
SEQ ID nº 4782	2646	467	SA-1340.2	1964390-1965202 p	similar to 5-keto-D-gluconate 5-reductase
SEQ ID nº 4783	2647	468	SA-1341.1	1965219-1965857 p	similar to sugar-phosphate isomerase
SEQ ID n° 4784	2648	469	SA-1342.1	1965883-1966890 p	similar to 2-keto-3-deoxygluconate kinase
SEQ ID n° 4785	- 2649	470	SA-1343.1	1,966902-1967540 p	similar to 2-dehydro-3-deoxyphosphogluconate aldolase-
SEQ ID n° 4786	2650	471	SA-1344.2	1968248-1970101 p	Unknown
SEQ ID n° 4787	2651	472	SA-1345.1	1970272-1970886 m	similar to unknown protein
SEQ ID n° 4788	2652	473	SA-1347.1	1971019-1971444 m	similar to transcription regulator (MarR family)
SEQ ID n° 4789	2653	474	SA-1348.1	1971571-1975977 m	DNA polymerase III (alpha subunit)
SEQ ID nº 4790	2654	475	SA-1349.1	1976100-1976684 m	similar to peptidoglycan hydrolase
SEQ ID n° 4791	2655	477	SA-1350.1	1976809-1978662 m	prolyl-tRNA synthetase
SEQ ID n° 4792	2656	478	SA-1351.1	1978754-1980013 m	similar to other protein
					similar to phosphatidate cytidylyltransferase (CDP-diglyceride
SEQ ID n° 4793	2657	479	SA-1352.1	1980044-1980838 m	synthase)
SEQ ID n° 4794	2658	480	SA-1353.3	1980853-1981605 m	similar to undecaprenyl pyrophosphate synthetase
SEQ ID n° 4795	2659	481	SA-1354.2	1218666-1219799 m	similaer to hypothetical permeases
SEQ ID n° 4796	2660	482	SA-1355.1	1219890-1221326 m	similar to 6-phospho-beta-glucosidase
SEQ ID n° 4797	2661	483	SA-1356.1	1221489-1222103 m	similar to other proteins
SEQ ID n° 4798	2662	484	SA-1357.1	1222204-1223025 m	similar to unknown protein
SEQ ID n° 4799	2663	485	SA-1358.1	1223650-1224579 p	similar to transcriptional regulator (AraC/XyIS family)
SEQ ID n° 4800	2664	486	SA-1359.1	1224636-1226168 m	similar to putative channel transporter
SEQ ID n° 4801	2665	487	SA-136.1	264314-265096 p	Unknown
					similar to spermidine/putrescine ABC transporter (binding
SEQ ID n° 4802	2666	488	SA-1361.3	1226277-1227350 m	protein)
SEQ ID n° 4803	2667	489	SA-1362.1	1227343-1228119 m	similar to spermidine/putrescine ABC transporter (permease)
SEQ ID n° 4804	2668	490	SA-1363.1	1228116-1228910 m	similar to spermidine/putrescine ABC transporter (permease)
					similar to spermidine/putrescine ABC transporter (ATP-binding
SEQ ID n° 4805	5669	491	SA-1364.1	1228894-1230048 m	protein)
SEQ ID n° 4806	2670	492	SA-1365.1	1230097-1230999 m	similar to UDP-N-acetylenolpyruvoylglucosamine reductase
SEQ ID n° 4807	2671	493	SA-1366.1	1231143-1231631 m	similar to hydroxymethylpterin pyrophosphokinase
SEQ ID n° 4808	2672	494	SA-1367.1	1231628-1231990 m	similar to dihydroneopterin aldolase
SEQ ID n° 4809	2673	495	SA-1368.1	1231992-1232795 m	similar to dihydropteroate synthase
SEQ ID n° 4810	2674	496	SA-137.1	265093-266226 m	similar to integrase
SEQ ID n° 4811	2675	497	SA-1370.1	1232799-1233362 m	similar to GTP cyclohydrolase



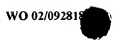
SEQ ID n° 4812	2676	498	SA-1371.1	1233381-1234643 m	Similar to folyl-polyglutamate synthetase
SEQ ID n° 4813	2677	499	SA-1372.1	1234645-1235532 m	similar to unknown protein
SEQ ID n° 4814	2678	200	SA-1373.2	1235519-1236385 m	similar to homoserine kinase
SEQ ID n° 4815	2679	501	SA-1374.3	1236387-1237670 m	similar to homoserine dehydrogenase
SEQ ID n° 4816	2680	502	SA-1377.1	1647764-1649380 m	Similar to nickel ABC transporter (binding protein)
SEQ ID n° 4817	2681	503	SA-1378.1	1646833-1647777 m	Similar to nickel ABC transporter (permease)
SEO 10 p. 4818	7687	504	SA_1370 1	1646027 1646833 m	Similar to constitution but to constitution of reliming
SEO ID nº 4819	2683	505	SA-138 1	266230-766436 m	similar to unknown proteins
SEQ ID n° 4820	2684	506	SA-1380.1	1645251-1646039 m	Similar to olioppentide ABC transporter (ATP-hinding protein)
					Similar to oligopeptide or nickel ABC transporter (ATP-binding
SEQ ID n° 4821	2685	507	SA-1381.1	1644584-1645264 m	protein)
SEQ ID n° 4822	2686	508	SA-1382.1	1643735-1644463 m	uridylate kinase
SEQ ID n° 4823	2687	509	SA-1383.1	1643162-1643719 m	ribosome recycling factor
SEQ ID n° 4824	2688	510	SA-1384.1	1642190-1643044 m	similar to unknown proteins
SEQ ID n° 4825	2689	511	SA-1385.1	1641543-1642052 m	similar to peptide methionine sulfoxide reductase
SEQ ID n° 4826	2690	512	SA-1386.1	1641331-1641546 m	
	••••				similar to myosin-crossreactive streptococcal antigen (unknown
SEQ ID n° 4827	2691	513	SA-1388.1	1639401-1641173 m	function)
					similar to phosphate starvation inducible protein, unknown
SEQ ID n° 4828	2692	514	SA-1389.1	1638291-1639277 m	function
SEQ ID n° 4829	2693	515	SA-1390.2	1637397-1638200 p	Unknown
SEQ ID n° 4830	2694	516	SA-1391.2	2028876-2030531 m	Isimilar to two-component sensor histidine kinase
SEQ ID n° 4831	2695	517	SA-1392.1	2030524-2031201 m	similar to two-component response regulator (PhoB)
SEQ ID n° 4832	2696	518	SA-1393.1	2031201-2031857 m	Similar to transcriptional regulator PhoU
SEQ ID n° 4833	2697	519	SA-1394.1	2031854-2032603 m	similar to phosphate ABC transporter (ATP-binding protein)
SEQ ID n° 4834	2698	520	SA-1395.1	2032596-2033474 m	similar to phosphate ABC transporter (permease)
SEQ ID n° 4835	2699	521	SA-1396.1	2033476-2034321 m	similar to phosphate ABC transporter (permease)
SEQ 1D n° 4836	2700	523	SA-1399.1	2034336-2035217 m	similar to phosphate ABC transporter (binding protein)
SEQ ID n° 4837	2701	524	SA-14.1	1035400-1035831 m	similar to single-strand binding protein
SEQ ID n° 4838	2702	. 525	SA-140.1	266442-267230 m	similar to replication protein (plasmid)
SEQ ID n° 4839	2703	526	SA-1400.1	2035418-2036005 m	Unknown
SEQ ID n° 4840	2704	527	SA-1401.1	2036002-2036742 m	Similar to unknown proteins
SEQ ID n° 4841	2705	528	SA-1403.1	2036742-2037695 m	Similar to ribosomal protein L11 methyltransferase
SEQ ID n° 4842	2706	529	SA-1404.1	2037692-2038150 m	Similar to unknown proteins
SEQ ID n° 4843	2707	530	SA-1405.1	2038274-2038990 p	Similar to transcriptional regulators

SEQ ID n° 4844	2708	531	SA-1406.1	2039029-2039499 m	Similar to unknown proteins
SEQ 1D n° 4845	2709	532	SA-1407.1	2039471-2039929 m	Similar to unknown proteins
SEQ ID n° 4846	2710	533	SA-1408.1	2039916-2040059 m	Unknown
SEQ_ID_n° 4847	2711	534	SA-141.1	267411-267725 m	Unknown
SEQ ID nº 4848	2712	535	SA-1410.1	2040065-2040535 m	Similar to unknown proteins
					similar to xylulose-5-phosphate/fructose-6-phosphate
SEQ-ID n°-4849	i 2713	536	SA-1413.1	1907193-1909571 m	phosphoketolase
SEQ 1D n° 4850	2714	537	SA-1414.1	1909661-1910752 m	Similar to unknown proteins
SEQ ID n° 4851	2715	538	SA-1415.1	1911069-1912748 m	Weakly similar to transcriptional regulator (antiterminator)
SEQ ID n° 4852	2716	539	SA-1416.2	1912814-1913575 m	similar to unknown proteins
SEQ ID n° 4853	2717	541	SA-1418.2	1913595-1915112 m	similar to L-xylulose kinase
SEQ ID n° 4854	2718	542	SA-1419.1	1915178-1916167 m	Unknown
SEQ ID n° 4855	2719	543	SA-1421.1	1916189-1917640 m	Similar to galactitol-specific PTS enzyme IIC
SEQ ID n° 4856	2720	544	SA-1424.1	1917738-1918694 m	similar to glycerate dehydrogenase
SEQ ID n° 4857	2721	545	SA-1425.2	1918713-1919732 m	Unknown
SEQ ID nº 4858	2722	546	SA-1427.1	154278-155117 p	similar to undecaprenol kinase
SEQ ID n° 4859	2723	547	SA-1429.1	155237-155992 p	similar to competence negative regulator mecA
SEQ ID n° 4860	2724	548	SA-143.1	267824-269182 m	weakly similar to DNA translocase
					similar to Streptococcus mutans RgpG protein required for
SEQ ID n° 4861	2725	549	SA-1430.1	155994-157154 p	biosynthesis of rhamnose-glucose polysaccharide
SEQ ID n° 4862	2726	220	SA-1431.1	157319-158089 p	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4863	2727	551	SA-1432.1	158126-159388 p	similar to unknown protein
SEQ ID n° 4864	2728	552	SA-1433.1	159390-160622 p	similar to aminotransferase, putative cysteine desulfurase
SEQ ID n° 4865	2729	553	SA-1434.1	160609-161052 p	similar to NifU protein
SEQ ID n° 4866	2730	554	SA-1435.1	161152-162570 p	similar to unknown protein
					similar to serine-type D-Ala-D-Ala carboxypeptidase (penicilin
SEQ ID n° 4867	2731	555	SA-1436.1	162642-163829 m	binding protein)
					similar to serine-type D-Ala-D-Ala carboxypeptidase (penicilin
SEQ ID n° 4868	2732	556	SA-1437.2	163982-165217 m	binding protein)
SEQ ID n° 4869	2733	557	SA-1438.2	613213-613473 m	50S ribosomal protein L31
SEQ ID n° 4870	2734	558	SA-1439.1	611580-613100 p	similar to metal ABC transporter (binding protein)
SEQ ID n° 4871	2735	559	SA-144.1	269145-269570 m	Unknown
SEQ ID n° 4872	2736	560	SA-1440.1	610041-611438 p	similar to dipeptidase
SEQ ID n° 4873	2737	561	SA-1441.1	609115-610026 p	Similar to unknown proteins
SEQ ID n° 4874	2738	562	SA-1442.2	608144-609118 p	Similar to unknown proteins
SEQ ID n° 4875	2739	563	SA-1445.2	607257-608147 p	Similar to unknown proteins

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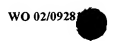
SEQ ID nº 4876	2740	564	SA-1446.1	606690-607103 m	Similar to unknown proteins
SEQ ID n° 4877	2741	565	SA-1447.1	605461-606444 m	Similar to purine nucleoside hydrolase
SEQ ID n° 4878	2742	566	SA-1448.1	604907-605464 m	Similar to unknown proteins
SEQ ID n° 4879	2743	567	SA-1449.1	603521-604867 p	asparaginyl-tRNA synthetase
SEQ ID nº 4880	2744	999	SA-145.1	269636-269917 m	similar to unknown proteins
SEQ ID n° 4881	2745	569	SA-1450.1	602307-603500 p	similar to aspartate aminotransferase
SEQ ID n° 4882	2746	570	SA-1451.1	599714-602221 p	similar to ATP-dependent DNA helicase
SEQ 10 n° 4883	2747	571	SA-1453.2	598842-599606 p	Similar to meso-2,3-butanediol dehydrogenase (D-acetoin forming)
SEQ 1D n° 4884	2748	573	SA-1457.2	1931763-1934015 p	similar to gamina-glutarnyicysteine synthetase (for the North Jacob Landinal part)
SEQ ID n° 4885	2749	574	SA-1458.1	1930919-1931593 p	Similar to unknown proteins
SEQ ID n° 4886	2750	575	SA-146.1	270084-270542 p	Unknown
SEQ ID n° 4887	2751	576	SA-1460.1	1929535-1930557 m	Similar to transcriptional regulators and to PTS enzyme II
SEQ ID n° 4888	2752	577	SA-1462.1	1927872-1929164 m	Similar to adenylosuccinate synthase
SEQ ID n° 4889	2753	578	SA-1463.1	1927556-1927849 p	Unknown
SEQ ID n° 4890	2754	579	SA-1465.1	1926600-1927247 m	Similar to unknown protein
SEQ ID n° 4891	2755	580	SA-1466.1	1924993-1926432 m	Putative PTS enzyme IIC
SEQ ID n° 4892	2756	581	SA-1467.1	1924687-1924965 m	similar to mannitol-specific PTS enzyme IIB
SEQ ID n° 4893	2757	582	SA-1468.1	1924135-1924620 m	similar to mannitol-specific PTS enzyme IIA
SEQ ID n° 4894	2758	583	SA-1469.1	1923357-1924022 m	Similar to hexulose-6-phosphate synthase
SEQ ID n° 4895	2759	584	SA-147.1	270919-271479 p	similar to putative acetyl transferase
SEQ ID n° 4896	2760	585	SA-1470.1	1922490-1923353 m	Similar to hexulose-6-phosphate isomerase
SEQ ID n° 4897	2761	586	SA-1471.1	1921772-1922488 m	Similar to L-ribulose-5-phosphate 4-epimerase
SEQ 1D n° 4898	2762	587	SA-1472.3	1921114-1921761 m	similar to hypothetical transaldolase
SEQ ID n° 4899	2763	589	SA-1474.1	104233-104421 p	Unknown
SEQ ID n° 4900	2764	590	SA-1475.1	104755-105075 p	Unknown
SEQ ID n° 4901	2765	591	SA-1477.1	105865-106416 p	Similar to unknown proteins
SEQ ID n° 4902	2766	592	SA-1478.1	106636-107055 p	Similar to unknown proteins
SEQ ID n° 4903	2767	593	SA-1479.1	107255-107734 p	Similar to the putative sigma factor ComX1
SEQ ID n° 4904	2768	594	SA-148.1	271579-272157 p	similar to putative acetyl transferase
SEQ ID n° 4905	2769	595	SA-1480.1	107856-108548 p	Similar to phosphoglycerate mutase
SEQ ID n° 4906	2770	596	SA-1481.1	108545-109297 p	Similar to D,D-carboxypeptidase
SEQ ID n° 4907	2771	597	SA-1482.1	109294-109869 p	Similar to N-acetyl muramidase



SEQ ID n° 4908	2772	598	SA-1483.1	110012-111046 p	Similar to transcription repressor of class I heat-shock (HrcA)
SEQ ID n° 4909	2773	599	SA-1484.1	111049-111621 p	Similar to heat shock protein GrpE
SEQ ID n° 4910	2774	900	SA-1486.3	111802-113631 p	Chaperone protein DnaK
SEQ ID n° 4911	2775	601	SA-1487.3	113920-115059 p	Chaperone protein DhaJ
SEQ ID n° 4912	2776	604	SA-149.1	272173-272880 p	similar to unknown protein
SEQ ID n° 4913	2777	605	SA-1490:2	1650563-1650988 m	ribosomal protein L11
SEQ ID n° 4914	2778	909	SA-1491.1	1651199-1652581 m	similar to transmembrane efflux pump protein
SEQ ID n° 4915	2779	209	SA-1492.1	1652589-1653803 m	similar to hypothetical N-acyl-L-amino acid amidohydrolase
SEQ ID n° 4916	2780	809	SA-1493.1	1654062-1654967 p	similar to transcriptional regulator (LysR family)
SEQ ID n° 4917	2781	609	SA-1494.1	1655028-1655381 p	similar to unknown proteins
SEQ ID n° 4918	2782	610	SA-1495.1	1655410-1657125 m	similar to para-aminobenzoate synthase, component I
SEQ ID n° 4919	2783	611	SA-1498.1	1657207-1659648 m	Similar to DNA translocase
SEQ ID n° 4920	2784	612	SA-1499.1	1659823-1660626 p	similar to peptidyl-prolyl cis-trans isomerase
					similar to surface antigen proteins, putative peptidoglycan bound
SEQ ID n° 4921	2785	613	SA-15.3	745991-748792 p	protein (LPXTG motif)
SEQ ID n° 4922	2786	614	SA-150.1	273140-274087 m	similar to unknown proteins
SEQ ID n° 4923	2787	615	SA-1500.1	1660678-1661511 m	similar to metal ABC transporter (permease)
SEQ ID n° 4924	2788	616	SA-1501.1	1661513-1662229 m	similar to metal ABC transporter (ATP-binding protein)
SEQ ID n° 4925	2789	617	SA-1502.2	1662400-1663326 m	similar to metal ABC transporter (binding protein)
					similar to C5A peptidase, putative peptidoglycan linked protein
SEQ ID n° 4926	2790	618	SA-1503.2	2080002-2084714 p	(LPXTG motif)
SEQ ID n° 4927	2791	619	SA-1504.1	2084947-2085633 p	similar to two-component response regulator
SEQ ID n° 4928	2792	620	SA-1506.1	2085630-2087018 p	similar to two-component sensor histidine kinase
SEQ ID n° 4929	2793	621	SA-1507.1	2087111-2087719 p	Unknown
SEQ ID n° 4930	2794	622	SA-151.1	274084-274575 m	similar to RNA polymerase ECF sigma factor
SEQ ID n° 4931	2795	623	SA-1511.2	2087806-2090307 m	leucyl-tRNA synthetase
SEQ ID n° 4932	2796	624	SA-1512.2	356072-356626 p	similar to unknown proteins
SEQ ID n° 4933	2797	625	SA-1513.1	355330-355995 p	similar to competence protein ComFC
SEQ ID n° 4934	2798	627	SA-1515.1	354041-355330 p	competence protein ComFA
SEQ ID n° 4935	2799	628	SA-1516.1	353341-353985 m	similar to unknown proteins
SEQ ID n° 4936	2800	629	SA-1517.1	352324-353250 m	similar to cysteine synthetase A
SEQ ID n° 4937	2801	630	SA-1518.1	351084-352172 p	similar to glycerol dehydrogenase
SEQ ID n° 4938	2802	631	SA-1519.1	350348-351016 p	similar to hypothetical transaldolase
SEQ ID n° 4939	2803	632	SA-152.1	275157-275765 m	similar to transcriptional regulator (TetR/AcrR family)
SEQ ID n° 4940	2804	633	SA-1520.3	347882-350338 p	similar to formate acetyltransferase



SEQ ID n° 4941	2805	634	SA-1521.1	1269394-1269624 m	similar to unknown proteins
SEQ ID n° 4942	2806	635	SA-1522.1	1269707-1270729 m	similar to branched-chain-amino-acid aminotransferase
SEQ ID n° 4943	2807	636	SA-1523.1	1270842-1273301 m	Similar to topoisomerase IV subunit A
SEQ ID n° 4944	2808	637	SA-1524.1	1273435-1275384 m	similar to DNA topoisomerase IV (subunit B)
SEQ ID n° 4945	2809	638	SA-1525.1	1275498-1276136 p	similar to unknown proteins
SEQ ID n° 4946	2810	639	SA-1526.1	1276202-1276855 m	Uracii DNA glycosylase
SEQ ID n° 4947	2811	640	SA-1527.1	1276954-1277439 m	similar to unknown proteins
SEQ ID n° 4948	2812	641	SA-1528.1	1277553-1278794 m	CMP-N-acetylneuraminic acid synthetase
SEQ ID n° 4949	2813	642	SA-153.1	275789-276886 m	Similar to ABC transporter (permease)
SEQ ID n° 4950	2814	643	SA-1530.1	1278805-1279434 m	similar to glycosylation or acetylation protein
SEQ ID n° 4951	2815	644	SA-1531.1	1279431-1280585 m	similar to UDP-N-acetylglucosamine-2-epimerase
SEQ ID n° 4952	2816	645	SA-1532.1	1280662-1281687 m	similar to N-acetylneuraminic acid synthetase
SEQ ID n° 4953	2817	646	SA-1533.1	1281687-1283087 m	capsular polysaccharide repeat unit transporter
SEQ ID n° 4954	2818	647	SA-1535.2	973749-974447 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4955	2819	648	SA-1536.1	972955-973737 m	putative ABC transporter (permease)
SEQ ID n° 4956	2820	648	SA-1537.1	972314-972919 p	similar to unknown transmembrane protein
SEQ ID n° 4957	2821	029	SA-1538.1	970960-971529 m	similar to acetyltransferase (chloramphenicol ?)
SEQ ID n° 4958	2822	651	SA-1539.1	969736-970839 p	Unknown
SEQ ID n° 4959	2823	653	SA-1543.1	966944-969736 p	similar to cation-transporting P-ATPase
SEQ ID n° 4960	2824	654	SA-1544.1	966358-966792 p	similar to peptide methionine sulfoxide reductase
SEQ ID n° 4961	2825	655	SA-1545.1	965894-966358 p	similar to unknown proteins
SEQ ID n° 4962	2826	929	SA-1546.1	965232-965885 p	similar to unknwon proteins
SEQ ID n° 4963	2827	299	SA-1547.2	962493-965126 p	weakly similar to histidine triad protein, putative lipoprotein
SEQ ID n° 4964	2828	658	SA-155.1	276889-277605 m	1.2 Transport/binding proteins and lipoproteins
SEQ ID n° 4965	2829	629	SA-1551.2	656055-658727 p	Putative peptidoglycan bound protein (FPXTG motif)
SEQ ID n° 4966	2830	099	SA-1552.1	658766-659008 p	similar to hypothetical sortase protein (N-terminal part)
SEQ ID n° 4967	2831	661	SA-1554.1	658966-659571 p	similar to putative surface protein (sortase)
SEQ ID n° 4968	2832	662	SA-1555.1	659787-660425 p	Unknown
SEQ ID n° 4969	2833	664	SA-1558.1	661523-661699 p	Unknown
₽	2834	665	SA-156.2	277869-278552 m	similar to unknown proteins
SEQ ID n° 4971	2835	999	SA-1560.1	663083-663313 p	Unknown
SEQ ID n° 4972	2836	299	SA-1562.1	663374-664525 p	Putative cell wall protein, weakly similar to peptidase or esterase
SEQ ID n° 4973	2837	899	SA-1563.1	664725-665717 p	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4974	2838	699	SA-1564.1	665720-666538 p	similar to unknown proteins, putative transmembrane protein
SEQ 1D n° 4975	2839	029	SA-1565.2	666540-667325 p	similar to unknown proteins, putative transmembrane protein



SEQ ID n° 4976	2840	671	SA-1567.3	1831820-1832131 m	Similar to unknown proteins
SEQ ID n° 4977	2841	672	SA-1568.3	1832242-1833135 p	Similar to Ribonuclease HII
SEQ ID n° 4978	2842	673	SA-157.2	278573-278884 m	similar to unknown proteins
₽	2843	674	- SA-1570.1	1833151-1833744 p	Similar to signal peptidase 1
SEQ ID n° 4980	2844	675	SA-1571.2	1833873-1836293 p	Similar to putative exodeoxyribonuclease V
SEQ ID n° 4981	2845	676	SA-1573.3	1836407-1836889 p	Similar to unknown proteins
SEQ ID n° 4982	2846	. 677	SA-1574.1	1836960-1838054 m	Similar to DNA-damage-inducible protein P
SEQ ID n° 4983	2847	678	SA-1577.1	1838239-1840551 p	Similar to Pyruvate formate-lyase
SEQ ID n° 4984	2848	629	SA-1578.1	1840652-1841026 p	Similar to FMN-binding protein
SEQ ID n° 4985	2849	680	SA-1579.1	1841061-1841990 m	Similar to unknown proteins
SEQ ID n° 4986	2850	681	SA-1580.1	1841987-1842742 m	Similar to C3-degrading proteinase
SEQ ID n° 4987	2851	682	SA-1582.1	1842867-1843763 p	Similar to unknown protein
SEQ ID n° 4988	2852	683	SA-1584.3	1843842-1844690 m	Similar to glycerol uptake facilitator
SEQ ID n° 4989	2853	684	SA-1585.3	1844859-1845311 m	Similar to unknown protein
SEQ ID n° 4990	2854	685	SA-1586.2	1845329-1846531 m	Similar to efflux protein
SEQ ID n° 4991	2855	686	SA-1587.2	624205-625581 p	similar to ABC transporter (permease)
SEQ ID n° 4992	2856	687	SA-1588.1	623549-624205 p	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 4993	2857	688	SA-1589.1	622262-623539 p	similar to ABC transporter (permease)
SEQ ID n° 4994	2858	689	SA-159.2	279076-279783 p	similar to other proteins
SEQ ID n° 4995	2859	690	SA-1590.1	621359-621520 m	similar to unknown proteins
SEQ ID n° 4996	2860	691	SA-1591.1	620914-621186 p	similar to transposase
SEQ ID n° 4997	2861	692	SA-1593.1	620591-620851 p	similar to transposase
SEQ ID n° 4998	2862	693	SA-1594.1	620369-620569 p	similar to transposase
SEQ ID n° 4999	2863	694	SA-1595.1	619803-620207 p	similar to insertion elemant protein
SEQ ID n° 5000	2864	695	SA-1596.1	618920-619540 m	similar to integrase C-terminal part (truncated)
SEQ ID n° 5001	2865	696	SA-1597.1	618428-618775 p	ribosomal protein L19
SEQ ID n° 5002	2866	697	SA-1598.1	616652-617848 p	similar to unknown proteins
SEQ ID n° 5003	2867	698	SA-1599.1	616384-616659 p	similar to other proteins
ů	2868	700	SA-160.2	280061-281209 p	similar to N-acetylglucosamine-6-phosphate deacetylase
SEQ ID n° 5005	2869	701	SA-1601.1	615864-616307 p	similar to flavodoxin
SEQ ID n° 5006	2870	702	SA-1602.1	614783-615805 p	similar to adenosine deaminase
SEQ ID n° 5007	2871	703	SA-1603.2	613582-614517 m	similar to unknown proteins
SEQ ID n° 5008	2872	704	SA-1604.2	1867026-1867295 p	30S ribosomal protein S14
SEQ ID n° 5009	2873	705	SA-1605.2	1867657-1868682 m	similar to low specificity L-threonine aldolase
SEO ID nº 5010	2874	706	SA-1606.2	1868802-1869812 m	Similar to other proteins (includin putative glycoprotein endopeptidase)
2122		2			

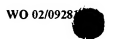


SEO ID nº 5011	2875	707	54-1607 1	1860888-1870334 m	Similar to putative acetyltransferase
					Similar to other proteins (includin putative disconposition
SEQ 1D n° 5012	2876	708	SA-1608.1	1870297-1870986 m	endopeptidase)
SEQ ID nº 5013	2877	709	SA-1609.1	1871168-1871398 p	Similar to unknown proteins
SEQ ID n° 5014	2878	710	SA-161.2	1457718-1458158 m	similar to hypothetical transcriptional regulator
SEQ 1D n° 5015	2879	711	SA-1610.1	1871452-1873131 p	Similar to unknown proteins
SEQ ID n° 5016	2880	712	SA-1611.1 · -·	1873293-1873802 m	Similar to secreted unknown proteins -
SEQ ID n° 5017	2881	713	SA-1612.1	1873950-1875296 m	glutamine synthetase .
SEQ ID n° 5018	2882	714	SA-1613.1	1875330-1875701 m	Similar to transcriptional regulator, MerR/GlnR family
SEQ ID n° 5019	2883	715	SA-1614.1	1875781-1876320 m	Similar to unknown protein
SEQ ID n° 5020	2884	717	SA-1616.1	1876583-1877779 m	Similar to phosphoglycerate kinase
SEQ ID n° 5021	2885	718	SA-1617.2	1877914-1878783 m	Similar to unknown lipoprotein
SEQ ID n° 5022	. 3886	719	SA-1619.2	2071134-2071673 p	similar to other proteins
SEQ ID n° 5023	2887	720	SA-162.1	1455854-1457671 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5024	2888	721	SA-1620.1	2071874-2072959 p	similar to glycerol dehydrogenase
SEQ ID n° 5025	2889	722	SA-1621.1	2073124-2073831 m	similar to unknown protein
SEQ ID n° 5026	2890	723	SA-1622.1	2074178-2076022 m	similar to other proteins
SEQ ID n° 5027	2891	724	SA-1623.1	2076067-2078304 m	similar to cobalamin-independent methionine synthase MetC
SEQ ID n° 5028	2892	725	SA-1624.1	2078674-2078997 m	similar to unknown protein
SEQ ID n° 5029	2893	726	SA-1625.2	2078987-2079679 m	similar to putative transport protein
SEQ ID n° 5030	2894	727	SA-1626.3	517632-518201 p	Similar to unknown proteins
SEQ ID n° 5031	2895	728	SA-1627.1	518299-518883 p	Similar to unknown proteins
SEQ ID n° 5032	2896	729	SA-1628.1	518880-519446 p	Similar to unknown proteins
SEQ ID n° 5033	, 2897	730	SA-1629.1	519446-522100 p	valyl-tRNA synthetase
SEQ ID n° 5034	2898	731	SA-163.1	1454110-1455864 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5035	2899	732	SA-1630.1	522336-523265 m	Similar to unknown proteins
SEQ ID n° 5036	2900	733	SA-1632.1	523682-524641 p	'Similar to oxidoreductase
SEQ ID n° 5037	2901	734	SA-1634.1	524802-525704 p	Similar to putative divalent cation transport protein
SEQ ID n° 5038	2902	735	SA-1635.3	525864-526928 p	similar to unknown proteins
SEQ ID n° 5039	2903	736	SA-1636.2	1746531-1747739 m	Similar to transport proteins
SEQ ID n° 5040	2904	737	SA-1638.1	1745814-1746362 m	Similar to unknown proteins
SEQ ID n° 5041	2905	738	SA-164.1	1453376-1454002 m	similar to other proteins
SEQ ID n° 5042	2906	739	SA-1640.2	1744121-1745797 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5043	2907	740	SA-1641.2	1743298-1744128 m	Similar to hypothetical ABC transporter (permease)
SEQ ID n° 5044	2908	741	SA-1642.1	1742592-1743263 p	Similar to unknown proteins
SEQ ID n° 5045	2909	742	SA-1643.1	1741200-1742579 p	similar to Na+-transporting ATP synthase

SEQ ID n° 5046	2910	743	SA-1644.1	1740472-1741185 p	Similar to glucose-inhibited division protein GidB
SEQ ID nº 5047	2911	744	SA-1645.1	1739790-1740344 m	Similar to unknown proteins
SEQ ID n° 5048	2912	745	SA-1646.1	1738879-1739769 m	Similar to other proteins
SEQ ID n° 5049	2913	746	SA-1647.1	1738241-1738774 m	Similar to unknown proteins
SEQ ID n° 5050	2914	747	SA-1648.1	1737317-1738006 m	Similar to two-component response regulator
SEQ ID n° 5051	2915	748	SA-1649.2	1735822-1737327 m	Similar to two-component sensor histidine kinase
SEQ ID n° 5052	2916	749	SA-165.1	1452688-1453374 m	Similar to two-component response regulator —
SEQ ID n° 5053	2917	750	SA-1650.2	1636729-1637205 m	similar to other proteins
SEQ ID n° 5054	2918	751	SA-1651.1	1635601-1636407 m	similar to unknown proteins
SEQ ID n° 5055	2919	752	SA-1653.1	1634709-1635206 m	similar to unknown proteins
SEQ ID n° 5056	2920	753	SA-1654.1	1634330-1634728 m	similar to diacylglycerol kinase
SEQ ID n° 5057	2921	754	SA-1655.1	1633389-1634288 m	similar to GTP binding proteins
SEQ ID n° 5058	2922	755	SA-1656.1	1632751-1633152 m	Unknown
SEQ ID n° 5059	2923	757	SA-1658.1	1631767-1632021 m	Unknown
SEQ ID n° 5060	2924	759	SA-166.1	1451459-1452688 m	Similar to two-component sensor histidine kinase
SEQ ID n° 5061	2925	160	SA-1660.1	1629232-1630824 p	ู่ปกหกอพท
SEQ ID n° 5062	2926	761	SA-1662.1	1628167-1629015 m	similar to transcriptional regulator
SEQ ID n° 5063	2927	762	SA-1663.1	1627197-1628018 m	similar to formamidopyrimidine-DNA glycosylase
SEQ ID n° 5064	2928	763	SA-1665.1	1626613-1627200 m	similar to dephosphocoenzyme A kinase
SEQ ID n° 5065	2929	764	SA-1667.1	1625788-1626489 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5066	2930	765	SA-1669.1	1624152-1625798 m	Unknown
SEQ ID n° 5067	2931	992	SA-167.1	1450465-1451343 m	similar to mevalonate kinase
SEQ ID n° 5068	2932	797	SA-1671.2	1622787-1623995 m	similar to efflux pump
SEQ ID n° 5069	2933	768	SA-1673.2	1988716-1990323 m	Similar to glucan 1,6-alpha-glucosidase
SEQ ID n° 5070	2934	769	SA-1674.1	1987633-1988628 m	Similar to UDP-galactose 4-epimerase
SEQ ID n° 5071	2935	770	SA-1675.1	1986932-1987612 p	similar to two-component response regulator
SEQ ID.n° 5072	2936	771	SA-1676.1	1985404-1986930 p	similar to two-component sensor histidine kinase
SEQ ID n° 5073	2937	772	SA-1677.1	1983913-1985250 m	similar to organic acid transport protein
SEQ ID n° 5074	2938	774	SA-1679.4	1982725-1983888 m	similar to malic enzyme ((S)-malate:NAD+ oxidoreductase)
SEQ ID n° 5075	2939	775	SA-168.1	1449539-1450483 m	similar to mevalonate diphosphate decarboxylase
SEQ 10 n° 5076	2940	776	SA-1680.2	1942117-1942581 m	Similar to transctiptional regulator CtsR
SEQ ID n° 5077	2941	777	SA-1681.2	1939673-1942120 m	Similar to endopeptidase Clp ATP-binding chaln C
SEQ ID n° 5078	2942	778	SA-1682.1	1938962-1939453 p	Similar to acyltransferase
SEQ ID n° 5079	2943	779	SA-1683.1	1938307-1938948 p	similar to deoxypurine kinase subunit
SEQ ID n° 5080	2944	780	SA-1685.1	1937209-1938186 p	Similar to transcription regulator
SEQ ID n° 5081	2945	781	SA-1686.1	1936350-1937225 p	Similar to other proteins



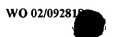
SEQ ID n° 5082	2946	782	SA-1687.1	1934954-1936210 p	Similar to other proteins
SEQ 1D n° 5083	2947	783	SA-1689.3	1934139-1934957 p	Similar to unknown proteins
SEQ 1D n° 5084	2948	784	SA-169.1	1448554-1449546 m	similar to phosphomevalonate kinase
SEQ ID n° 5085	2949	786	SA-1694.1	28975-30318 p	cell wall separation
SEQ ID n° 5086	2950	788	SA-1696.1	31518-32693 p	similar to aminotransferase
SEQ ID n° 5087	2951	789	SA-1697.2	32683-33444 p	similar to unknown protein
SEQ ID n° 5088	2952	790	SA-1698.2	1725796-1726299 m	Similar to unknown proteins
SEQ ID n° 5089	2953	791	SA-1699.1	1726385-1727716 m	Similar to UDP-N-acetylmuramate-alanine ligase
SEO ID nº 5090	2954	792	\$4-17.1	1038856-1030578 m	similar to unknown protein, putative peptidoglycan bound protein
SEQ ID n° 5091	2955	793	SA-170.1	1447562-1448557 m	similar to Isopentenyl diphosphate isomerase
SEQ ID n° 5092	2956	794	SA-1700.1	1727726-1728343 m	Similar to unknown proteins
SEQ ID n° 5093	2957	795	SA-1702.1	1728508-1731606 m	Similar to SWI/SNF family helicase
SEQ ID n° 5094	2958	796	SA-1704.1	1731762-1733072 m	Similar to GTP binding protein
SEQ ID n° 5095	2959	797	SA-1705.1	1733120-1734022 m	Similar to primosome component (helicase loader) Dnal
SEQ ID n° 5096	, 2960	798	SA-1706.2	1734019-1735194 m	Similar to unknown proteins
SEQ ID n° 5097	2961	199	SA-1707.2	1735194-1735673 m	Similar to unknown protein
SEQ ID n° 5098	2962	800	SA-171.1	1447276-1447494 m	Unknown
SEQ ID n° 5099	2963	801	SA-1710.2	2102209-2103849 m	similar to unknown protein
SEQ ID n° 5100	2964	802	SA-1711.1	2101372-2102184 p	similar to other protein
SEQ ID n° 5101	2965	803	SA-1712.1	2098935-2101256 m	similar to penicillin-binding protein 2a
SEQ ID n° 5102	2966	804	SA-1714.1	2098525-2098698 m	similar to other protein
SEQ ID n° 5103	2967	806	SA-1716.2	2096282-2098213 m	putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 5104	2968	807	SA-1718.2	1809587-1810966 m	similar to ABC transporter (ATP-binding protein)
					Similar to unknown protein, putative ABC transporter
SEQ ID n° 5105	2969	808	SA-1719.1	1810959-1811672 m	(permease)
SEQ ID n° 5106	2970	808	SA-172.1	1446376-1447230 p	similar to unknown proteins
ଠା	2971	810	SA-1720.1	1811672-1812268 m	Similar to unknown protein
°c	2972	811	SA-1722.1	1812279-1812710 m	Similar to unknown protein
SEQ ID n° 5109	2973	812	SA-1723.1	1812725-1812880 m	Unknown
•					Similar to 2,3-dihydroxybenzoate-AMP ligase (C-terminal
SEQ ID n° 5110	2974	813	SA-1724.1	1812894-1813433 m	domain)
0.00	1	3	1		Similar to 2,3-dihydroxybenzoate-AMP ligase (enterobactin
SEQ 10 n° 3111	CJ67	814	SA-1/25.1	1813437-1814261 m	synthetase component E)
SEQ 1D n° 5112	2976	815	SA-1726.1	1814469-1814843 m	Similar to transcription regulator



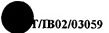
SEQ ID n° 5113	2977	816	SA-1728.1	1814891-1815343 m	Similar to late competence protein required for DNA binding and ubtake comEB
SEQ ID n° 5114	2978	817	SA-1729.1	1815355-1816422 m	Similar to similar to Xaa-Pro dipeptidase
SEQ ID n° 5115	2979	818	SA-173.1	1445361-1446275 p	similar to unknown proteins
SEQ ID n° 5116	2980	819	SA-1730.1	1816522-1817238 m	Similar to unknown protein
SEQ ID n° 5117	2981	820	SA-1731.2	1817240-1818739 m	similar to multidrug resistance protein
۵۳	2982	821	SA-1732.2	959863-960279 p	nucleoside-diphosphate kinase
SEQ ID n° 5119	2983	822	SA-1734.1	959293-959553 p	hypothetical
SEQ ID n° 5120	2984	824	SA-1737.1	957581-958246 p	similar to unknown proteins
SEQ ID n° 5121	2985	825	SA-1738.1	957253-957594 p	similar to unknown proteins
SEQ ID n° 5122	2986	826	SA-1739.1	956387-957256 p	similar to unknown proteins
SEQ ID n° 5123	2987	827	SA-174.1	1444618-1445274 p	similar to other proteins
SEQ ID n° 5124	2988	828	SA-1740.2	952252-956385 p	similar to unknown proteins
SEQ ID n° 5125	2989	829	SA-1741.2	2040862-2041353 p	Similar to putative acetyl transferase
SEQ ID n° 5126	2990	830	SA-1742.1	2041346-2042614 p	Similar to unknown proteins
SEQ ID n° 5127	2991	831	SA-1744.1	2043157-2043462 m	Unknown
SEQ ID n° 5128	2992	832	SA-1745.1	2043446-2043847 m	Unknown
SEQ ID n° 5129	2993	833	SA-1746.1	2043835-2045043 m	Similar to unknown proteins
SEQ ID n° 5130	2994	834	SA-1747.1	2045045-2045437 m	Unknown
SEQ ID n° 5131	2995	835	SA-1748.1	2045504-2045959 m	weakly similar to integrase
SEQ ID n° 5132	2996	837	SA-175.1	1444176-1444625 p	similar to unknown proteins
SEQ ID n° 5133	2997	838	SA-1750.1	2046386-2046622 m	hypothetical gene
SEQ ID n° 5134	2998	839	SA-1751.1	2046794-2047288 p	Similar to transcriptional regulator (phage related)
SEQ 10 n° 5135	2999	840	SA-1753.2	774016-774840 p	Similar to unknown proteins
의	3000	841	SA-1754.1	773219-774016 p	Similar to unknown proteins
اءْ	3001	842	SA-1755.1	769589-773128 p	Similar to chromosome segregation SMC protein
٥٥	3002	843	SA-1756.1	768895-769581 p	Similar to ribonuclease III
SEQ ID n° 5139	3003	844	SA-1757.1	768351-768719 p	Similar to unknown proteins
SEQ ID n° 5140	3004	845	SA-1758.1	767539-768348 p	Similar to unknown proteins
SEQ ID n° 5141	3005	846	SA-1759.1	766186-767535 p	Similar to two-component sensor histidine kinase
SEQ ID n° 5142	3006	847	SA-176.1	1442801-1444084 p	similar to 3-hydroxy-3-methylglutaryl-coenzyme A reductase
SEQ 1D n° 5143	3007	848	SA-1760.1	765483-766193 p	similar to two-component response regulator
SEQ ID n° 5144	3008	850	SA-1762.2	1503364-1503699 m	Similar to unknown proteins
SEQ ID n° 5145	3009	851	SA-1763.1	1503783-1504949 m	Similar to chorismate synthase
SEQ ID n° 5146	3010	852	SA-1764.1	1504950-1506017 m	Similar to 3-dehydroquinate synthase
SEQ ID n° 5147	3011	853	SA-1765.1	1506111-1506788 m	similar to 3-dehydroquinate dehydratase



SEQ ID n° 5148	3012	854	SA-1766.1	1506788-1507945 m	Similar to unknown proteins
SEQ ID n° 5149	3013	855	SA-1767.1	1508080-1510224 p	Similar to unknown proteins
ID n°	3014	856	SA-1768.2	1510465-1510824 m	50S ribosomal protein L20
SEQ ID n° 5151	3015	857	SA-1769.2	1513077-1513274 p	Similar to ferredoxin
					similar to 3-hydroxy-3-methylglutaryl-coenzyme A synthase
	3016	858	SA-177.1	1441627-1442799 p	(HMG-CoA synthase)
SEQ ID n° 5153	3017	-859	SA-1770.1	1513261-1513752 m	Similar to unknown protein
SEQ ID n° 5154	3018	098	SA-1771.1	, 1513781-1515001 m	Similar to tripeptidase
SEQ ID n° 5155	3019	861	SA-1772.1	1515137-1516771 m	Similar to unknown protein
					Similar to UDP-N-acetylmuramoylalanyl-D-glutamyl-2, 6-
SEQ ID n° 5156	3020	862	SA-1773.1	1516895-1518349 p	diaminopimelate ligase
SEQ ID n° 5157	3021	863	, SA-1774.2	1518504-1519298 p	similar to ferrichrome ABC transporter (ATP-binding protein)
SEQ ID n° 5158	3022	864	SA-1775.2	1519322-1520254 p	similar to ferrichrome ABC transporter (binding protein)
SEQ ID n° 5159	3023	985	SA-1776.1	1520270-1521295 p	similar to ferrichrome ABC transporter (permease)
ID n°	3024	998	SA-1778.1	1521292-1522293 p	Similar to ferrichrome ABC transporter (permease)
SEQ ID n° 5161	3025	298	SA-1779.2	1522323-1522976 m	Similar to unknown protein
SEQ ID n° 5162	3026	898	SA-178.1	1440634-1441473 m	thymidylate synthase
SEQ ID n° 5163	3027	698	SA-1780.2	327377-328462 p	Similar to two-component sensor histidine kinase
SEQ ID n° 5164	3028	870	SA-1781.1	326640-327380 p	Similar to ABC transporter (permease)
υ υ	3029	871	SA-1783.2	325757-326653 p	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5166	3030	872	SA-1784.2	324021-325628 p	similar to unknown proteins
° □	3031	873	SA-1785.1	323350-323832 m	similar to autoinducer-2 production protein (LuxS)
SEQ ID n° 5168	3032	874	SA-1787.1	321662-323125 p	similar to unknown proteins
SEQ ID n° 5169	3033	875	SA-1788.1	320495-321649 p	similar to unknown proteins
SEQ ID n° 5170	3034	876	SA-179.1	1440060-1440554 m	similar to dihydrofolate reductase
SEQ ID n° 5171	3035	877	SA-1791.2	319693-320025 p	similar to unknown proteins
SEQ ID n° 5172	3036	878	SA-1792.2	1469012-1470220 m	similar to poly(A) polymerase
SEQ ID n° 5173	3037	879	SA-1793.1	1467133-1469001 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5174	3038	880	SA-1794.1	1466663-1467136 m	similar to unknown proteins
SEQ ID n° 5175	3039	881	SA-1795.1	1464857-1466596 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5176	3040	882	SA-1796.1	1463083-1464852 m	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5177	3041	883	SA-1797.2	1462533-1463042 p	similar to unknown proteins
o u	3042	884	SA-1798.3	1568308-1569804 m	Similar to 4-alpha-glucanotransferase (amylomaltase)
SEQ ID n° 5179	3043	885	SA-1799.3	1566032-1568296 m	Similar to glycogen phosphorylase
SEO ID p. 5180	3077	900	0 0 0	40006EF 4044040	similar to plasmid surface exclusion protein, putative
טוני די טוני	2014	000	JA-10.1	1038655-1041916 m	peptidoglycan bound protein (LPA1G motif)

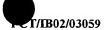


SEQ ID n° 5248	3112	958	SA-188.1	1434490-1435866 m	similar to amino acid transporter
SEQ ID n° 5249	3113	929	SA-1881.2	82132-82437 p	ribosomal protein L24
SEQ ID n° 5250	3114	960	SA-1882.2	82461-83003 p	ribosomal protein L5
SEQ ID n° 5251	3115	961	SA-1883.1	83361-83759 p	ribosomal protein S8
SEQ ID n° 5252	3116	962	SA-1884.1	83869-84405 p	ribosomal protein L6
SEQ ID n° 5253	3117	963	SA-1885.1	84506-84862 p	ribosomal protein L/18
SEQ ID n° 5254	3118	964	SA-1886.1	84881-85375 p	ribosomal protein S5
SEQ ID n° 5255	3119	996	SA-1888.1	85390-85569 p	ribosomal protein L30
SEQ ID n° 5256	3120	296	SA-1891.1	85694-86134 p	ribosomal protein L15
SEQ ID n° 5257	3121	998	SA-1892.1	86155-87459 p	similar to preprotein translocase SecY
SEQ ID n° 5258	3122	696	SA-1893.2	87554-88192 p	adenylate kinase
SEQ ID n° 5259	3123	970	SA-1894.2	61615-62718 p	Unknown
SEQ ID n° 5260	3124	971	SA-1897.1	62804-64102 p	similar to adenylosuccinate lyase
SEQ ID n° 5261	3125	972	SA-1898.1	64256-65158 p	similar to unknown proteins
'n	3126	973	SA-1899.1	65447-66445 p	similar to Holliday junction DNA helicase, subunit B
ື	3127	974	SA-190.1	1433541-1434485 m	similar to homocysteine S-methyltransferase
SEQ ID n° 5264	3128	975	SA-1900.3	66597-67034 p	similar to protein-tyrosine phosphatase
SEQ ID n° 5265	3129	926	SA-1901.2	974448-974816 m	similar to transcriptional regulator (GntR family)
SEQ ID n° 5266	3130	977	SA-1902.2	974961-978065 p	DNA polymerase III (alpha subunit)
SEQ ID n° 5267	3131	978	SA-1904.1	978146-979168 p	similar to 6-phosphofructokinase
SEQ ID n° 5268	3132	626	SA-1906.1	979217-980719 p	similar to pyruvate kinase
SEQ ID n° 5269	3133	980	SA-1908.2	980890-981447 p	similar to type-1 signal peptidase
SEQ ID n° 5270	3134	981	SA-1909.2	1996039-1996464 m	Similar to galactose 6-P isomerase (A subunit)
					similar to ATP-dependent Clp proteinase (ATP-binding subunit),
SEQ ID n° 5271	3135	982	SA-191.2	1431339-1433447 p	ClpL
SEQ ID n° 5272	3136	. 983	SA-1910.1	1995503-1996018 m	Similar to galactose 6-P isomerase (B subunit)
SEQ ID n° 5273	3137	984	SA-1911.1	1994560-1995492 m	Similar to tagatose-6-phosphate kinase
ů	3138	985	SA-1912.1	1993581-1994558 m	Similar to tagatose 1,6-diP aldolase
SEQ ID n° 5275	3139	986	SA-1913.1	1992631-1993527 m	Similar to unknown proteins
SEQ ID n° 5276	3140	286	SA-1915.1	1991685-1992536 m	Similar to unknown proteins
SEQ ID n° 5277	3141	988	SA-1916.2	1990451-1991584 m	Similar to sugar ABC transporter (ATP-binding protein)
SEQ ID n° 5278	3142	989	SA-1918.3	2104019-2105230 m	similar to phosphopentomutase
SEQ ID n° 5279	3143	066	SA-1919.1	2105297-2105968 m	similar to deoxyribose-phosphate aldolase
SEQ ID n° 5280	3144	991	SA-1921.1	2105998-2107200 m	similar to transport system permease protein
SEQ ID n° 5281	3145	992	SA-1922.1	2107221-2108000 m	similar to uridine phosphorylase
SEQ ID n° 5282	3146	993	SA-1923.1	2108158-2108895 p	similar to transcriptional regulator (GntR family)



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SEQ ID n° 5283	314/	994	SA-1924.1	2108908-2109204 p	Unknown
SEQ 1D n° 5284	3148	966	SA-1926.2	2109304-2110926 m	chaperonin GroEL
SEQ ID n° 5285	3149	666	SA-1929.2	1348211-1349551 m	similar to multidrug resistance protein
SEQ ID n° 5286	3150	1000	SA-193.2	1430405-1430905 m	Ribosomal protein L10
SEQ ID n° 5287	3151	1001	SA-1930.1	1347385-1348152 p	Similar to unknown proteins
SEQ ID n° 5288	3152	1002	SA-1931.1	1345445-1347226 m	excinuclease ABC (subunit C)
					Transmembrane N-terminal domain, C-terminal domain similar
SEQ ID n° 5289	3153	1003	SA-1932.2	1343622-1345403 p	to hydrolases
SEQ ID n° 5290	3154	1004	SA-1933.2	1342850-1343452 m	similar to NADH dehydrogenase
SEQ ID n° 5291	3155	1005	SA-1934.2	1341397-1342803 m	similar to dipeptidase
SEQ ID n° 5292	3156	1006	SA-1935.1	1340716-1341300 m	similar to unknown proteins
SEQ ID n° 5293	3157	1001	SA-1937.1	1339967-1340701 m	similar to unknown proteins
SEQ ID n° 5294	3158	1009	SA-194.1	1429976-1430341 m	ribosomal protein L7/L12
SEQ ID n° 5295	3159	1010	SA-1940.3	782771-783706 p	Hpr (ser) kinase/phosphatase
SEQ ID n° 5296	3160	1012	SA-1942.1	782280-782543 p	Similar to unknown proteins
SEQ ID n° 5297	3161	1013	SA-1943.1	781741-782199 p	Similar to unknown proteins
SEQ ID n° 5298	3162	1014	SA-1944.1	779616-781778 p	Similar to unknown proteins
SEQ ID n° 5299	3163	1015	SA-1945.1	778468-779511 p	Similar to oxidoreductase
SEQ ID n° 5300	3164	1016	SA-1946.1	778207-778335 m	Similar to unknown proteins
SEQ ID n° 5301	3165	1017	SA-1949.1	777299-778201 m	Similar to unknown proteins
SEQ ID n° 5302	3166	1020	SA-1951.1	776487-777299 m	Similar to unknown proteins
					Similar to signal recognition particle and to cell division protein
SEQ ID n° 5303	3167	1021	SA-1952.2	774840-776450 p	FtsY
					similar to glycine betaine/carnitine/choline ABC transporter (ATP-
SEQ ID n° 5304	3168	1022	SA-1953.2	256648-257793 m	binding protein)
SEQ ID n° 5305	3169	1023	SA-1954.1	256013-256648 m	similar to choline ABC transporter (permease)
1	,				similar to glycine betaine/carnitine/choline ABC transporter
SEQ ID n° 5306	3170	1024	SA-1955.1	255084-256010 m	(osmoprotectant-binding protein)
	,				similar to glycine betaine/carnitine/choline ABC transporter
SEQ ID n° 5307	3171	1025	SA-1956.1	254433-255083 m	(permease)
SEQ ID n° 5308	3172	1026	SA-1957.1	253190-254176 p	similar to efflux protein (truncated)
SEQ ID n° 5309	3173	1027	SA-1958.1	252257-253117 p	similar to transcriptional regulator (Rgg like)
SEQ ID n° 5310	3174	1028	SA-1959.1	251133-251414 p	Unknown
SEQ ID n° 5311	3175	1029	SA-196.1	, 1429562-1429735 m	Unknown
SEQ ID n° 5312	3176	1030	SA-1960.1	250713-251099 p	Unknown
SEQ ID n° 5313	3177	1031	SA-1961.1	250324-250728 p	bseudogene

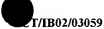
SEQ ID n° 5385	3249	1109	SA-2056.1	844892-846148 p	Similar to permease (gluconate ?)
SEQ ID n° 5386	3250	1110	SA-2059.2	843725-844867 p	Similar to putative glycerate kinase
SEQ ID n° 5387	3251	1111	SA-206.1	1425120-1425608 m	similar to unknown proteins
SEQ ID n° 5388	3252	1112	SA-2060.1	842464-843558 m	Similar to unknown proteins
SEQ ID n° 5389	3253	1113	SA-2061.2	840968-842395 p	Similar to beta-glucosidase
SEQ ID n° 5390	3254	1114	SA-2062.2	81399-81659 p	ribosomal protein S17
SEQ ID n° 5391	3255	1115	SA-2063.2	81167-81373 p	ribosomal protein L29
SEQ ID n° 5392	3256	1116	SA-2065.1	80744-81157 p	ribosomal protein L16
SEQ ID n° 5393	3257	1117	SA-2066.1	80087-80740 p	ribosomal protein S3
SEQ ID n° 5394	3258	1118	SA-2067.1	79730-80074 p	ribosomal protein L22
SEQ ID n° 5395	3259	1119	SA-2069.1	79436-79714 p	ribosomal protein S19
SEQ ID n° 5396	3260	1120	SA-207.1	1423303-1425120 m	similar to plasmid transfer complex protein TrsK
SEQ ID n° 5397	3261	1121	SA-2071.1	78504-79337 p	ribosomal protein L2
SEQ ID n° 5398	3262	1122	SA-2072.1	78190-78486 p	ribosomal protein L23
SEQ ID n° 5399	3263	1123	SA-2073.1	77567-78190 p	ribosomal protein L4
SEQ ID n° 5400	3264	1124	SA-2074.2	76917-77543 p	ribosomal protein L3
SEQ ID nº 5401	3265	1125	SA-2075.2	76504-76812 p	ribosomal protein S10
SEQ ID n° 5402	3266	1127	SA-2077.3	1351826-1352680 p	similar to transposase (truncated)
SEQ ID n° 5403	3267	1128	SA-2078.2	1363404-1363694 p	similar to transposase, N-terminal part
SEQ ID n° 5404	3268	1129	SA-2079.2	.1362139-1363350 m	Unknown
SEQ ID n° 5405	3269	1130	SA-208.1	1423041-1423283 m	Unknown
SEQ ID n° 5406	3270	1131	SA-2082.3	1358151-1361603 m	streptococcal C5a peptidase
SEQ ID n° 5407	3271	1132	SA-2083.4	1800964-1802403 p	Similar to sucrose-6-phosphate hydrolase
SEQ ID n° 5408	3272	1133	SA-2084.1	1802405-1803367 p	Similar to transcriptional regulator (Laci family)
SEQ ID n° 5409	3273	1134	SA-2085.1	1803454-1803888 m	Similar to transcription termination protein NusB
SEQ ID n° 5410	3274	1135	SA-2086.1	1803881-1804270 m	Similar to unknown proteins
SEQ ID n° 5411	3275	1136	SA-2087.1	1804359-1804919 m	Similar to translation elongation factor EF-P
SEQ ID n° 5412	3276	1137	SA-2090.1	1806098-1807819 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5413	3277	1139	SA-2092.2	1807809-1809566 m	similar to ABC transporter (binding protein)
SEQ ID n° 5414	3278	1140	SA-2095.2	464590-465939 p	glucose-6-phosphate isomerase
SEQ ID n° 5415	3279	1141	SA-2096.1	466261-466788 p	similar to unknown protein
SEQ ID n° 5416	3280	1142	SA-2097.1	466785-467456 p	similar to unknown protein
SEQ ID n° 5417	3281	1143	SA-2099.1	467588-468631 p	similar to putative ABC transporter (binding protein)
SEQ ID n° 5418	3282	1144	SA-21.1	1043024-1043371 m	Unknown
SEQ ID n° 5419	3283	1145	SA-210.1	1422170-1423024 m	similar to unknown protein from conjugative transposon
SEQ ID n° 5420	3284	1146	SA-2100.1	468722-469621 m	similar to UDP-glucose pyrophosphorylase



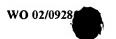
SEQ ID n° 5421	3285	1147	SA-2101.1	469658-470674 m	similar to glycerol-3-phosphate dehydrogenase
°	3286	1148	SA-2102.1	470844-471173 p	similar to ribonuclease P protein component
°c	3287	1149	SA-2103.2	471186-472001 p	similar to unknown protein and to B. subtilis SpolliJ protein
°c	3288	1150	SA-2104.2	2047351-2048703 m	Unknown
SEQ ID n° 5425	3289	1151	SA-2105.1	2048713-2049804 m	Unknown
SEQ ID n° 5426	3290	1152	SA-2106.1	2050044-2050988 m	similar to hypothetical transcriptional regulator
SEQ ID n° 5427	3291	1153	SA-2107.1	2051052-2051441 m	Unknown
SEQ ID n° 5428	3292	1154	SA-2109.1	2052439-2052756 p	Unknown
SEQ ID n° 5429	3293	1155	SA-211.1	1421755-1422108 m	similar to unknown protein from conjugative transposons
SEQ ID n° 5430	3294	1156	SA-2110.1	2052805-2052933 p	Unknown
SEQ ID n° 5431	3295	1157	SA-2112.1	2053374-2053664 p	Similar to unknown proteins
SEQ ID n° 5432	3296	1158	SA-2113.1	2053763-2054137 p	Unknown
SEQ ID n° 5433	3297	1159	SA-2115.1	2054427-2055155 p	Unknown
SEQ ID n° 5434	3298	1160	SA-2116.1	2055450-2055644 p	Unknown
SEQ ID n° 5435	3299	1161	SA-2117.1	2055763-2056128 m	Unknown
SEQ ID n° 5436	3300	1162	SA-2118.2	2056356-2056754 m	Unknown
SEQ ID n° 5437	3301	1163	SA-2119.2	1523036-1523971 m	Similar to manganese-dependent inorganic pyrophosphatase
SEQ ID n° 5438	3302	1164	SA-212.1	1419453-1421798 m	similar to plamsid transfer complex protein TraE
SEQ ID n° 5439	3303	1165	SA-2120.1	1524088-1524876 m	Similar to pyruvate-formate lyase activating enzyme
SEQ ID n° 5440	3304	1166	SA-2122.1	1524944-1526278 m	Similar to unknown protein
°	3305	1167	SA-2123.1	1526458-1527024 m	Similar to unknown proteins
SEQ ID n° 5442	3306	1168	SA-2124.1	1527017-1527952 m	Similar to unknown protein
SEQ ID n° 5443	1 3307	1169	SA-2125.3	1528045-1528686 m	Similar to unknown protein
SEQ ID n° 5444	3308	1170	SA-2126.3	1528667-1529251 m	similar to unknown protein
SEQ ID n° 5445	3309	1171	SA-2128.2	1352904-1353194 p	similar to transposase
SEQ ID n° 5446	3310	1172	SA-2129.1	1353352-1353732 p	degenerate transposase
SEQ ID n° 5447	3311	1173	SA-2130.1	1353776-1354036 p	degenerate trnasposase
SEQ ID n° 5448	3312	1174	SA-2131.1	1354082-1354315 p	Degenerate transposase
SEQ ID n° 5449	3313	1175	SA-2132.1	1354514-1356982 m	Similar to histidine triad protein
SEQ ID n° 5450	3314	1176	SA-2133.2	1356995-1357915 m	laminin-binding surface protein
SEQ ID n° 5451	3315	1177	SA-2134.2	625678-626331 p	similar to two-component response regulator VncR
SEQ ID n° 5452	3316	1178	SA-2136.1	626328-627647 p	similar to two-component sensor histidine kinase VncS
SEQ ID n° 5453	3317	1179	SA-2137.1	627699-628358 m	Similar to transposase C-terminal end (truncated)
SEQ ID n° 5454	3318	1180	SA-2138.1	628524-628724 p	Similar to unknown proteins
SEQ ID n° 5455	3319	1182	SA-214.1	1416656-1419451 m	similar to plasmid and conjugative transposon protein
SEQ ID n° 5456	3320	1183	SA-2140.i	629379-630584 p	similar to FtsW and RodA proteins

SEQ ID n° 5457	3321	1184	SA-2141.1	630703-631263 p	similar to unknown protein
SEQ ID n° 5458	3322	1185	SA-2143.2	631264-633216 p	DNA gyrase, subunit B
SEQ ID n° 5459	3323	1186	SA-2145.1	1900897-1901571 m	similar to two-component response regulator
SEQ ID n° 5460	3324	1187	SA-2146.1	1901927-1902061 m	ribosomal protein L34
SEQ ID n° 5461	3325	1188	SA-2147.1	1902245-1903600 m	Similar to unknown proteins
SEQ 1D n° 5462	3326	1189	SA-2148.1	1903847-1905574 m	similar to betaine ABC transporter permease and substrate binding protein
0.00	1000	7.400	00770	0700007	
SEC 10 n 5463	3327	1190	SA-2149.2	1905593-1906816 m	similar to giycine betaine ABC transporter (ATP-binding protein)
SEQ ID n° 5464	3328	1191	SA-2156.2	2095537-2096076 m	similar to transcription antitermination factor nusG
SEQ ID n° 5466	3330	1193	SA-2158.1	2093024-2094220 p	similar to alvoosyl transferase
SEQ ID n° 5467	3331	1194	SA-2159.1	2091974-2092819 m	similar to unknown protein
SEQ ID n° 5468	3332	1195	SA-216.1	1415704-1416552 p	similar to unknown proteins
SEQ ID n° 5469	3333	1196	SA-2160.2	2090684-2091931 p	similar to transporter
SEQ ID n° 5470	3334	1197	SA-2161.2	1244116-1245459 p	similar to staphylokinase and streptokinase
SEQ ID n° 5471	3335	1198	SA-2162.1	1243011-1243697 m	similar to unknown protein
SEQ ID n° 5472	3336	1199	SA-2163.1	1241896-1242903 m	similar to unknown protein
SEQ ID n° 5473	3337	1200	SA-2165.1	1240446-1241822 m	similar to succinic semialdehyde dehydrogenase
SEQ ID n° 5474	3338	1201	SA-2166.1	1239758-1240303 m	similar to glycine betaine transporter (N-terminal end)
SEQ ID n° 5475	3339	1202	SA-2167.1	1238756-1239757 m	similar to glycine betaine transporter (C-terminal end)
SEQ ID n° 5476	3340	1203	SA-2168.1	1238248-1238733 p	similar to unknown protein C-terminal part
SEQ ID n° 5477	3341	1204	SA-2169.3	1237847-1238407 p	similar to unknown protein (N-terminal part)
SEQ ID n° 5478	3342	1205	SA-217.1	1415117-1415707 p	similar to unknown proteins
SEQ ID n° 5479	3343	1207	SA-2172.2	1100661-1102145 m	similar to carbon starvation protein A
SEQ ID n° 5480	3344	1208	SA-2173.1	1102301-1103035 m	similar to two-component response regulator lytR
SEQ ID.n° 5481	3345	1209	SA-2174.1	1103047-1104786 m	similar to two-component sensor histidine kinase LytS
SEQ ID n° 5482	3346	1212	SA-2178.1	1106331-1106708 m	Unknown
SEQ ID n° 5483	3347	1213	SA-2180.3	1107097-1107420 m	Unknown
SEQ ID n° 5484	3348	1214	SA-2182.3	1107768-1108301 m	Unknown
SEQ ID n° 5485	3349	1215	SA-2184.1	825634-827220 p	Similar to ATP-dependent RNA helicase, DEAD-box family
SEQ ID n° 5486	3350	1216	SA-2185.1	824569-825399 p	putative ABC transporter (binding protein)
SEQ ID n° 5487	3351	1217	SA-2186.1	823891-824553 p	similar to ABC transporter (permease)
SEQ ID n° 5488	3352	1218	SA-2187.1	823164-823898 p	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5489	3353	1219	SA-2188.1	822663-823043 p	Similar to unknown proteins
SEQ ID n° 5490	3354	1220	SA-2190.1	821033-822577 p	Similar to peptide-chain-release factor 3

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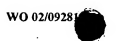
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SEO ID n° 5491	3355	1221	SA-2192 2	819318-820856 p	Olffillal to sufface proteins, purative peptidogrycan illined protein
SEQ ID nº 5492	3356	1222	SA-2195.3	1140455-1142050 m	similar to cardiolipin synthetase
SEQ ID n° 5493	3357	1223	SA-2196.2	1142169-1143839 m	Similar to formate-tetrahydrofolate ligase
SEQ ID n° 5494	3358	1224	SA-2197.2	1143928-1144947 m	similar to lipoate-protein ligase
SEQ ID n° 5495	3359	1225	SA-2198.2	1144974-1145852 m	similar to unknown protein
			-		similar to ATP-dependent Clp protease ATP-binding subunit
SEQ ID n° 5496	3360	1226	SA-22.1	1043463-1045295 m	ClpA
SEQ ID n° 5497	3361	1227	SA-220.1	1410000-1414904 m	Putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 5498	3362	1228	SA-2200.2	120068-121351 p	similar to trigger factor (prolyl isomerase)
SEQ ID n° 5499	3363	1229	SA-2201.1	119069-119911 m	similar to unknown protein
SEQ ID n° 5500	3364	1230	SA-2202.1	118463-119032 p	similar to unknown protein
SEQ ID n° 5501	3365	1231	SA-2203.1	118002-118466 p	similar to unknown protein
SEQ ID n° 5502	3366	1232	SA-2204.1	117234-117992 p	similar to phosphomethylpyrimidine kinase
SEQ ID n° 5503	3367	1233	SA-2205.1	116495-117271 p	similar to tRNA pseudouridine synthase A
SEQ ID n° 5504	3368	1234	SA-2206.2	115177-116424 p	Similar to other proteins
SEQ 1D n° 5505	3369	1235	SA-2207.2	573586-574413 p	similar to unknown proteins
SEQ ID n° 5506	3370	1236	SA-2208.2	574400-574873 p	similar to transcriptional regulator
SEQ ID n° 5507	3371	1237	SA-2210.1	574885-576543 p	similar to DNA repair and recombination protein RecN
SEQ ID n° 5508	3372	1238	SA-2212.1	576656-577492 p	similar to unknown proteins
SEQ ID n° 5509	3373	1239	SA-2213.1	577467-578324 p	similar to unknown proteins
SEQ ID n° 5510	3374	1240	SA-2214.2	578299-578901 p	Similar to unknown proteins
SEQ ID n° 5511	3375	1242	SA-2216.2	1250993-1251574 m	similar to unknown proteins
SEQ ID n° 5512	3376	1243	SA-2217.1	1250789-1250983 m	similar to unknown protein
SEQ 1D n° 5513	3377	1244	SA-2219.1	1250191-1250733 m	similar to unknown protein
SEQ ID n° 5514	3378	1245	SA-222.1	1409808-1409999 m	Unknown
SEQ ID n° 5515	3379	1246	SA-2220.1	1249935-1250132 m	similar to unknown protein
SEQ ID n° 5516	3380	1247	SA-2221.1	1249362-1249913 m	similar to unknown protein
SEQ ID n° 5517	3381	1248	SA-2222.1	1248557-1249198 m	similar to transcriptional regulator, GntR family
SEQ ID n° 5518	3382	1249	SA-2223.1	1247788-1248552 m	similar to unknown protein
SEQ ID n° 5519	3383	1250	SA-2224.1	1247129-1247788 m	similar to other protein
SEQ ID n° 5520	3384	1251	SA-2225.1	1246562-1247056 m	similar to thiol peroxidase
SEQ ID n° 5521	3385	1252	SA-2226.2	1245953-1246486 p	similar to unknown proteins
SEQ ID n° 5522	3386	1253	SA-2227.2	2001195-2001659 m	Similar to PTS enzyme IIA
SEQ ID n° 5523	3387	1254	SA-2228.1	2000888-2001193 m	Similar to PTS enzyme IIB
SEQ ID n° 5524	3388	1255	SA-2231.1	1999400-2000848 m	Similar to galactitol-specific PTS enzyme IIC



SEQ ID n° 5525	3389	1256	SA-2232.2	1996712-1999162 m	Similar to neuraminidase
SEQ ID n° 5526	3390	1257	SA-2233.2	1544787-1545722 m	Similar to putative rhamnosyltransferase
SEQ ID n° 5527	3391	1258	SA-2235.1	1545724-1546779 m	Similar to nucleptide-sugar dehydratase
SEQ ID n° 5528	3392	1259	SA-2236.1	1546782-1547501 m	Similar to 4-diphosphocytidyl-2C-methyl-D-erythritol synthase
000000000000000000000000000000000000000	0000	0007	7 1000 40	7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7 7	Similar to Pneumococcal LicD2 protein involved in
= °c	3394	1261	SA-2238 1	1547504-1548328 m 1548353-1550086 m	prosprovjenoline metabolism possible surface protein
SEQ ID n° 5531	3395	1262	SA-2239.1	1550079-1550432 m	Similar to unknown protein
SEQ ID n° 5532	3396	1263	SA-224.1	1409273-1409824 m	Unknown
SEQ ID n° 5533	3397	1264	SA-2240.1	1550429-1551157 m	Similar to putative glycosyl transferase
SEQ ID n° 5534	3398	1265	SA-2241.4	1551162-1552103 m	similar to rhamnosyltransferase
의	3399	1266	SA-2242.3	145779-146144 p	similar to unknown protein
SEQ ID n° 5536	3400	1267	SA-2244.3	146144-147808 p	similar to unknown protein
SEQ ID n° 5537	3401	1268	SA-2245.1	147956-148840 p	similar to unknown protein
SEQ ID n° 5538	3402	1270	SA-2247.1	149922-150662 m	similar to amind acid ABC transporter (ATP-binding protein)
					similar to amino acid (glutamine) ABC transporter (binding
SEQ 10 n° 5539	3403	1271	SA-2248.3	150672-152222 m	protein)
SEQ ID n° 5540.	3404	1272	SA-225.1	1408629-1409222 m	Unknown
SEQ ID n° 5541	3405	1275	SA-2253.4	1076528-1077445 m	similar to phosphate ABC transporter (permease)
SEQ ID n° 5542	3406	1276	SA-2254.2	1075651-1076538 m	similar to phosphate ABC transporter (permease)
SEQ ID n° 5543	3407	1277	SA-2255.1	1074836-1075639 m	similar to phosphate ABC transporter (ATP-binding protein)
SEQ ID n° 5544	3408	1278	SA-2256.1	1074066-1074824 m	similar to phosphate ABC transporter (ATP-binding protein)
SEQ ID n° 5545	3409	1279	SA-2258.2	1073379-1074032 m	similar to phosphate uptake regulatory protein
SEQ ID n° 5546	3410	1280	SA-2259.2	1086749-1089379 m	similar to hypothetical ABC transporter (permease)
SEQ ID n° 5547	3411	1281	SA-2260.1	1089391-1090092 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5548	3412	1282	SA-2261.3	1090229-1092349 m	similar to DNA thpoisomerase I
SEQ ID n° 5549	3413	1284	SA-2264.3	176720-177571 p	isimilar to 4-diphosphocytidyl-2-C-methyl-D-erythritol kinase
SEQ ID n° 5550	3414	1285	SA-2265.2	177657-178100 p	similar to transcriptional regulator
SEQ ID n° 5551	3415	1286	SA-2266.2	178103-178813 p	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5552	3416	1287	SA-2267.1	178803-179615 p	Similar to (metal) ABC transporter (permease)
SEQ ID n° 5553	3417	1288	SA-2268.2	179827-180858 m	similar to DNAse (competence related)
اء°	3418	1289	SA-2269.2	1617901-1619289 p	
SEQ ID n° 5555	3419	1290	SA-2271.1	1619373-1619840 m	Similar to small protein SmpB
°	3420	1291	SA-2272.2	1619843-1622248 m	similar to exoribbnuclease R
SEQ ID n° 5557	3421	1292	SA-2274.2	2111481-2112284 m	'Similar to ABC transporter (ATP-binding protein)



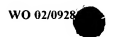
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Similar to ABC transporter (permease)	putative ABC transporter (binding protein)	Similar to unknown proteins	similar to unknown proteins	similar to unknown proteins	Unknown	Unknown	similar to signal recognition particle chain Fifth	Similar to unknown protein	similar to two-component sensor histidine kinase	Similar to unknown proteins	Similar to other proteins	Similar to competence protein ComEA	similar to competence protein CelB (ComEC)	Similar to unknown proteins	Similar to rhamnosyl transferase I	Similar to putative hexosyltransferase	Similar to transcription regulator	similar to plasmid DNA topolsomerase	similar to putative helicase and methylase	Similar to fibronectin binding protein, peptidoglycan bound	protein (LPXTG motif)	similar to ferrichrome ABC transporter (permease)	similar to (iron?) ABC transporter (permease)	similar to sugar O-acetyltransferase	similar to ribonuclease HII	similar to unknown protein, hypothetical GTP binding protein	similar to unknown proteins	similar to multidrug resistance protein	similar to hypothetical transcription factor	Unknown	group B oligopeptidase PepB	similar to hypothetical phosphoglycolate phosphatase	similar to methyltransferase	similar to protease maturation protein
2112289-2113185 m	2113201-2114163 m	2114628-2115425 m	2115611-2116471 p	2116514-2117245 p	1065700-1066194 m	1066231-1066419 m	1066541-1068106 m	1068124-1068456 m	1068545-1069858 m	827664-828428 m	828563-829303 p	829403-830056 p	830049-832277 p	832403-833212 p	1540196-1541044 m	1539067-1540206 m	1537486-1539015 p	1045297-1047045 m	1402425-1408625 m		1534497-1537202 m	1095272-1096246 m	1096243-1097205 m	1097444-1097992 m	1098013-1098774 m	1098761-1099612 m	1099888-1100460 m	852521-853672 p	853724-854671 p	1402054-1402353 m	854687-856492 p	856687-857313 p	857387-858094 p	858155-859084 p
SA-2276.1	SA-2278.1	SA-2279.1	SA-2280.1	SA-2281.2	SA-2282.2	SA-2283.1	SA-2285.1	SA-2287.1	SA-2288.2	SA-2289.2	SA-2291.1	SA-2292.1	SA-2294.2	SA-2295.2	SA-2296.2	SA-2297.1	SA-2298.1	SA-23.1	SA-230.1		SA-2300.3	SA-2302.2	SA-2303.1	SA-2304.1	SA-2305.1	SA-2306.1	SA-2307.2	SA-2308.2	SA-2309.1	SA-231.1	SA-2311.1	SA-2312.1	SA-2313.1	SA-2314.2
1293	1294	1295	1296	1297	1298	1299	1300	1301	1302	1303	1304	1305	1307	1308	1309	1310	1311	1312	1313		1314	1315	1316	1317	1318	1319	1320	1321	1322	1323	1324	1325	1326	1327
3422	3423	3424	3425	3426	3427	3428	3429	3430	3431	3432	3433	3434	3435	3436	3437	3438	3439	3440	3441		3442	3443	3444	3445	3446	3447	3448	3449	3450	3451	3452	3453	3454	3455
SEQ ID n° 5558	SEQ ID n° 5559	SEQ ID n° 5560	SEQ ID n° 5561	SEQ ID n° 5562	SEQ ID n° 5563	SEQ ID n° 5564	SEQ ID n° 5565	SEQ ID n° 5566	SEQ 1D n° 5567	SEQ ID n° 5568	SEQ ID n° 5569	SEQ ID n° 5570	SEQ ID n° 5571	SEQ ID n° 5572	SEQ ID n° 5573	SEQ ID n° 5574	SEQ ID n° 5575	SEQ ID n° 5576	SEQ ID n° 5577		SEQ ID n° 5578	SEQ ID n° 5579	SEQ ID n° 5580	SEQ ID n° 5581	SEQ ID n° 5582	SEQ ID n° 5583	SEQ ID n° 5584	SEQ ID n° 5585	SEQ ID n° 5586	SEQ ID n° 5587	SEQ ID n° 5588	SEQ ID n° 5589	SEQ ID n° 5590	SEQ ID n° 5591



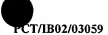
SEO ID nº 5592	3456	1328	CA_2345.2	211006E 21221E3 m	similar to anaerobic (class III) ribonucleotide reductase large
: °c	3457	1329	SA-2316 1	2119737-2119880 m	Jaknown
SEQ ID n° 5594	3458	1330	SA-2317 1	2118792-2119724 m	similar to oxidoreductase
SEQ ID n° 5595	3459	1331	SA-2318.1	2118292-2118783 m	similar to unknown proteins
ID n°	3460	1332	SA-2319.2	2117602-2118219 m	similar to anaerobic ribonucleotide reductase activator
SEQ ID n° 5597	3461	1333	SA-232.1	1401744-1402043 m	Unknown
ID nº	3462	1334	SA-2322.2	1501840-1503189 m	Similar to glutathione reductase
SEQ ID n° 5599	3463	1335	SA-2324.1	1501345-1501797 p	Similar to unknown protein (secreted protein)
SEQ ID n° 5600	3464	1336	SA-2326.1	1499953-1501098 m	Similar to iron-sulfur cofactor synthesis protein (NifS like)
SEQ ID n° 5601	3465	1337	SA-2327.2	1498737-1499951 m	Similar to hypothetical thiamine biosynthesis protein Thil
SEQ ID n° 5602	3466	1338	·SA-2329.3	1497457-1498635 m	similar to capsular polyglutamate biosynthesis
SEQ ID n° 5603	3467	1339	SA-233.1	1400941-1401579 m	Unknown
SEQ ID n° 5604	3468	1340	SA-2330.2	143337-144218 p	similar to fructose-bisphosphate aldolase class-II
SEQ ID n° 5605	3469	1341	SA-2331.2	144303-145220 m	similar to L-2-hydroxyisocaproate dehydrogenase
SEQ ID n° 5606	3470	1342	SA-2332.2	145459-145647 p	50S ribosomal protein L28
SEQ ID n° 5607	3471	1343	SA-2334.1	1532350-1534374 m	putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 5608	3472	1344	SA-2335.2	1531360-1532238 m	Similar to sortase protein
SEQ ID n° 5609	3473	1345	SA-2336.1	1530476-1531360 m	Similar to sortase protein
SEQ ID n° 5610	3474	1346	SA-2337.2	1529553-1530479 m	Putative peptidoglycan linked protein (LPXTG motif)
SEQ ID n° 5611	3475	1347	SA-2339.2	1289559-1290257 m	capsular polysaccharide chain length regulator/exporter
SEQ ID n° 5612	3476	1348	SA-234.1	1399815-1400900 m	similar to unknown proteins
SEQ ID n° 5613	3477	1349	SA-2340.2	1288158-1289546 m	similar to glucose-1-phosphate transferase
SEQ ID n° 5614	3478	1350	SA-2341.1	1287685-1288134 m	beta-1,4-galactosyltransferase enhancer
SEQ ID n° 5615	3479	1351	SA-2342.1	1287212-1287685 m	beta-1,4-galactosyltransferase
SEQ ID n° 5616	3480	1352	SA-2343.1	1286070-1287215 m	capsular polysaccharide repeating-unit polymerase
SEQ ID n° 5617	3481	1353	SA-2344.1	1285105-1286073 m	similar to glycosyl transferase
SEQ ID n° 5618	3482	1354	SA-2345.1	1284124-1285071 m	similar to glycosyl transferase
SEQ ID n° 5619	3483	1355	SA-2346.2	1283084-1284040 m	capsular polysaccharide biosynthesis protein
SEQ ID n° 5620	3484	1357	SA-2348.1	1217890-1218570 m	similar to DNA repair protein RadC
SEQ ID n° 5621	3485	1358	SA-2349.1	1217239-1217877 p	similar to unknown protein
°	3486	1359	SA-235.1	1399530-1399760 m	Unknown
SEQ ID n° 5623	3487	1360	SA-2350.1	1216737-1217084 p	similar to unknown protein
SEO ID 5° 5624	2700	1361	0 4 3 2 5 4 3	424EE20 424E73E x	similar to iron-sulfur cofactor synthesis protein or cysteine
4200 II OI	0010	100	2.1.62-00	J 5020-121-0205121	deaulius de la company de la c
SEQ ID n. 5625	3489	1364	SA-2354.1	1552093-1553247 M	similar to putative mamnosyitransierase



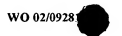
SEO 10 nº 5626 3490	1365	SA-2355 1	1553364-1554218 m	dTDP-L-rhamnose synthase
3491	1366	SA-2356.1	1554308-1554649 m	similar to unknown protein
3492	1367	SA-2357.1	1554758-1555867 m	RNA polymerase major sigma factor RpoD
3493	1368	SA-2358.2	1555875-1557683 m	DNA primase
3494	1369	SA-2359.3	121540-122115 p	similar to DNA-directed RNA polymerase (delta subunit)
3495	1370	SA-236.1	1399144-1399533 m	Unknown
3496	1371	SA-2360:1	122388-123992 p	similar to CTP synthetase
3497	1372	SA-2361.3	124101-125027 p	similar to unknown protein
3498	1374	SA-2364.2	1085765-1086643 m	similar to unknown proteins
3499	1375	SA-2365.1	1085068-1085658 m	Unknown
3500	1376	SA-2366.1	1083763-1085034 m	similar to unknown protein
3501	1377	SA-2367.1	1083319-1083750 m	similar to unknown protein
3502	1378	SA-2368.1	1082343-1083227 m	similar to tRNA pseudouridine 55 synthase
3503	1379	SA-2369.2	1081398-1082330 m	macrolide-efflux protein
3504	1380	SA-237.1	1398807-1399097 m	Unknown
3505	1381	SA-2370.3	1723907-1725709 m	Similar to other proteins
3506	1382	SA-2371.1	1723352-1723834 m	Similar to transcription elongation factor GreA
3507	1383	SA-2372.2	1721788-1723248 m	Similar to putative amidase
3508	1384	SA-2373.2	1560344-1561387 m	similar to ammonium transporter
3209	1385	SA-2374.1	1561467-1562594 p	Similar to unknown protein
3510	1386	SA-2376.1	1562631-1563617 m	Similar to unknown protein
3511	1387	SA-2377.1	1563780-1564085 p	Similar to unknown protein
3512	1388	SA-2378.2	1564243-1565616 m	Similar to glycerol (sugar)-3-phosphate transporter
3513	1389	SA-238.1	1398261-1398737 m	similar to phage repressor-like protein
3514	1390	SA-2380.2	890576-891346 p	similar to hydroxyethyl thiazole kinase (ThiM)
3515	1391	SA-2381.1	889777-890574 p	similar to phosphomethylpyrimidine kinase (ThiD)
3516	1392	SA-2382.1	889096-889752 p	Similar to putative thiamin biosynthesis protein
3517	1394	SA-2384.2	888320-889012 p	Similar to unknown proteins
3518	1395	SA-2386.1	1541041-1542465 m	Similar to capsular polysaccharide synthesis protein
3519	1396	SA-2387.1	1542465-1543829 m	Similar to unknown proteins
3520	1397	SA-2388.2	1543831-1544778 m	Similar to putative rhamnosyltransferase
3521	1398	SA-239.1	1397491-1398261 m	similar to unknown proteins
				similar to 5 -nucleotidase, putative peptidoglycan bound protein
3522	1399	SA-2390.2	1458427-1460499 p	(LPXTN motif)
3523	1400	SA-2391.1	1460536-1460946 m	similar to polypeptide deformylase
3524	1401	SA-2392.2	1461016-1462365 m	similar to NADP-specific glutamate dehydrogenase



SEQ ID n° 5661	3525	1402	] SA-2394.2	1133801-1135033 m	Similar to unknown protein
SEQ ID n° 5662	3526	1403	SA-2395.2	1135073-1136614 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5663	3527	1404	SA-2396.2	1480876-1481946 m	similar to other protein, putative transmembrane protein
°c	3528	1405	SA-2397.1	1482083-1483075 m	similar to thioredoxin reductase
SEQ ID n° 5665	3529	1406	SA-2398.1	1483056-1483808 m	similar to tRNA (quanine-N1)-methyltransferase
°	3530	1407	SA-2399.3	1483798-1484316 m	similar to 16S rRNA processing protein RimM
SEQ ID n° 5667	3531	1408	SA-240.2	1395238-1397478 m	similar to unknown proteins
°=	3532	1409	SA-2404.2	1798860-1800779 m	Similar to sucrose-specific PTS enzyme IIABC
	3533	1410	SA-2405.2	515623-516378 m	Similar to methyltransferase
SEQ ID n° 5670	3534	1411	SA-2406.1	515264-515602 m	similar to unknown proteins
SEQ ID n° 5671	3535	1412	SA-2407.1	514957-515262 m	similar to unknown proteins
SEQ ID n° 5672	3536	1413	SA-2409.2	514161-514739 p	Isimilar to integrase (C-terminal part)
의	3537	1414	SA-241.2	i 319054-319572 p	similar to unknown proteins
SEQ ID n° 5674	3538	1415	SA-2410.1	513604-514038 p	similar to integrase (N-terminal part)
°=	3539	1416	SA-2411.1	513228-513554 p	Similar to unknown phage proteins
SEQ ID n° 5676	3540	1417	SA-2412.1	512516-513055 p	Unknown
7F33 % CI O33					Similar to plasmid related proteins, Putative peptidoglycan bound
SEC 10 1 26/7	3541	1418	SA-2414.1	511516-512277 p	protein (LPXTG motif)
SEQ ID n° 5678	3542	1419	SA-2415.2	510934-511491 p	Unknown
SEQ ID n° 5679	3543	1420	SA-2416.2	510521-510934 p	Unknown
SEQ ID n° 5680	3544	1421	SA-2418.2	1069842-1070522 m	similar to two-component response regulator
SEQ ID n° 5681	3545	1422	SA-2419.2	1070684-1073233 m	similar to lysyl-aminopeptidase
SEQ ID n° 5682	3546	1423	SA-242.2	318380-318979 m	similar to recombination protein U (RecU)
SEQ ID n° 5683	3547	1424	SA-2420.2	1496953-1497267 m	50S ribosomal protein L21
SEQ ID n° 5684	3548	1426	SA-2422.2	1496608-1496946 m	Similar to unknown proteins
SEQ ID n° 5685	3549	1427	SA-2423.1	1496293-1496586 m	50S ribosomal protein L27
SEQ 10 n° 5686	3550	1428	SA-2424.1	1495172-1496077 m	Similar to transcriptional regulator (LysR/MarR family)
SEQ ID n° 5687	3551	1429	SA-2425.1	1494699-1495163 m	Similar to prolipoprotein signal peptidase
					Similar to ribosomal large subunit pseudouridine synthase B
SEQ ID n° 5688	3552	1430	SA-2426.3	1493825-1494715 m	(hypothetical)
SEQ ID n° 5689	3553	1431	SA-2428.3	1880206-1882284 m	translation elongation factor G
SEQ ID n° 5690	3554	1432	SA-2429.2	1265526-1266473 p	similar to unknown proteins
SEQ ID n° 5691	3555	1433	SA-2430.1	1266592-1267668 p	similar to bacteriophage integrase
SEQ ID n° 5692	3556	1434	SA-2431.3	1267848-1269050 m	similar to ribosomal protein S1-like DNA-binding protein
SEQ ID n° 5693	3557	1436	SA-2433.2	1942754-1943215 p	Similar to other proteins
SEQ ID n° 5694	3558	1437	SA-2434.4	1943374-1944414 m	translation elongation factor EF-Ts



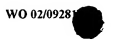
SEQ ID n° 5695	3559	1438	SA-2435.4	1944508-1945278 m	ribosomal protein S2
SEQ ID n° 5696	3560	1440	SA-2437.2	1290268-1290960 m	putative chain length regulator CpsC
SEQ ID n° 5697	3561	1441	SA-2438.1	1290969-1291700 m	Unknown
SEQ ID n° 5698	3562	1443	SA-244.2	316147-318384 m	similar to penicillin-binding protein 1A
SEQ ID n° 5699	3563	1444	SA-2440.3	1291706-1293163 m	Unknown
SEQ ID n° 5700	3564	1446	SA-2442.1	2153854-2154309 p	similar to other proteins (C-terminal end)
SEQ ID n° 5701	3565	1447	SA-2443.1	2152902-2153897 p	Unknown
SEQ ID n° 5702	3566	1448	SA-2445.2	2151206-2152471 p	similar to integrase
SEQ ID n° 5703	3567	1449	SA-2446.2	1369720-1370286 m	similar to plasmid replication protein
SEQ ID n° 5704	3568	1450	SA-2447.1	1369483-1369686 m	Unknown
SEQ ID n° 5705	3569	1451	SA-2448.1	1368284-1369465 m	Similar to integrase (phage-related protein)
SEQ ID n° 5706	3570	1452	SA-2450.2	1206681-1207373 p	Similar to unknown protein
SEQ ID n° 5707	3571	1454	SA-2453.2	849904-850605 p	similar to N-acetylglucosamine-6-phosphate isomerase
SEQ ID n° 5708	3572	1455	SA-2454.1	850677-851633 m	similar to unknown protein
SEQ ID n° 5709	3573	1456	SA-2455.2	851729-852448 p	similar to 16S pseudouridylate synthase
SEQ ID n° 5710	3574	1457	SA-2457.3	67418-69196 p	similar to unknown proteins
SEQ ID n° 5711	3575	1458	SA-2458.3	67041-67421 p	similar to unknown proteins
SEQ ID n° 5712	3576	1459	SA-2459.3	1715970-1716491 m	Similar to unknown proteins
SEQ ID n° 5713	3577	1460	SA-246.1	314767-316101 p	similar to cysteine aminopeptidase C
SEQ ID n° 5714	3578	1461	SA-2460.1	1716559-1717248 m	Similar to unknown proteins
SEQ ID n° 5715	3579	1462	SA-2461.1	1717271-1717774 m	Similar to unknown proteins
SEQ ID n° 5716	3580	1463	SA-2462.1	1717812-1718555 m	Similar to hypothetical rRNA methylase
SEQ ID n° 5717	3581	1464	SA-2463.1	1718587-1718865 p	similar to acylphosphatase
SEQ ID n° 5718	3582	1465	SA-2464.2	1718950-1719882 p	Similar to unknown proteins
SEQ ID n° 5719	3583	1467	SA-2466.2	715787-716482 m	Similar to amino acid ABC transporter (permease)
SEQ ID n° 5720	3584	1468	SA-2467.1	715125-715775 m	similar to ABC transporter (permease)
SEQ ID n° 5721	3585	1469	SA-2468.2	714510-715076 p	similar to unknown proteins
SEQ ID n° 5722	3586	1470	SA-2469.2	713324-714343 p	similar to unknown proteins
SEQ ID n° 5723	3587	1471	SA-247.1	313833-314654 p	similar to NH3-dependent NAD+ synthetase
SEQ ID n° 5724	3588	1472	SA-2470.2	785354-785629 m	Similar to unknown proteins
SEQ ID n° 5725	3589	1473	SA-2471.1	784882-785313 p	Similar to unknown proteins
SEQ ID n° 5726	3590	1474	SA-2472.1	784487-784885 p	Similar to unknown proteins
SEQ ID n° 5727	3591	1475	SA-2473.3	783699-784472 p	similar to prolipoprotein diacylglycerol transferase
SEQ ID n° 5728	3592	1479	SA-2477.2	1492009-1493079 m	Similar to carbamoyl-phosphate synthase, small subunit
SEQ ID n° 5729	3593	1480	SA-2479.2	1490355-1491953 m	similar to carbamoyl-phosphate synthase
SEQ ID n° 5730	3594	1481	SA-248.1	312376-313836 p	similar to unknown proteins



	2282	1482	SA-2480.2	1094514-1095275 m	similar to ferrichrome ABC transporter (ATP-binding protein)
(C)	3596	1483	SA-2482.2	1093424-1094452 m	similar to ferrichrome ABC transporter (binding protein)
3	3597	1484	SA-2483.2	1092444-1093286 m	similar to DNA processing Smf protein
3	3598	1485	SA-2486.2	1614454-1615296 m	similar to oxidoreductase
3	3599	1486	SA-2488.3	1488909-1490153 m	similar to unknown protein
3	3600	1487	SA-249.1	311304-312218 p	similar to thioredoxin reductase
3	3601	1488	SA-2490.2	1488196-1488906 m	similar to ABC transporter (ATP-binding protein)
Ē	3602	1489	SA-2491.2	1486934-1488181 m	similar to unknown protein.
ਲ 	3603	1490	SA-2492.2	981705-983519 p	similar to glucosamine-fructose-6-phosphate aminotransferase
ñ	3604	1491	SA-2493.2	1137253-1138329 m	similar to aspartate-semialdehyde dehydrogenase
Ř	3605	1493	SA-2495.1	1138538-1139770 m	highly repetitive peptidoglycan bound protein (LPXTG motif)
<u>~</u>	3606	1494	SA-2497.2	1080942-1081355 m	similar to unknown protein
Ñ	3607	1495	SA-2498.2	1080668-1080949 m	similar to unknown protein
3	3608	1496	SA-2499.3	1079914-1080678 m	similar to myo-inositol monophosphatase
Ř	3609	1497	SA-25.1	1047228-1051958 m	similar to plasmid proteins
36	3610	1498	SA-250.1	311011-311235 p	similar to unknwon proteins
36	3611	1499	SA-2500.3	1078536-1079846 m	conserved protein .
e E	3612	1500	SA-2501.3	1704700-1706052 m	similar to amino acid permease
3	3613	1501	SA-2502.2	1703979-1704527 m	Similar to rRNA methylase
3	3614	1502	SA-2503.2	1702674-1703843 m	similar to aminotransferase
Ř	3615	1503	SA-2504.2	1701935-1702564 m	uracil phosphoribosyltransferase
Э.	3616	1505	SA-2507.2	35466-35705 p	similar to acyl carrier protein
ř	3617	1506	SA-2508.1	34463-35455 p	similar to fatty acid/phospholipid synthesis protein
3	3618	1507	SA-2509.2	33507-34385 p	similar to unknown transmembrane protein
3(	3619	1508	SA-251.1	310146-310889 p	Similar to amino acid ABC transporter (ATP-binding protein)
36	3620	1509	SA-2510.2	1207444-1209444 m	similar to cation (K+) transport protein
3(	3621	1510	SA-2511.1	1209579-1210340 m	Similar to oxidoreductase
3(	3622	1511	SA-2512.2	1210402-1211394 m	phosphotransacetylase
36	3623	1512	SA-2513.2	329975-330604 p	similar to guanylate kinase
36	3624	1513	SA-2514.1	329104-329808 p	similar to unknown proteins
3	3625	1514	SA-2515.2	328464-329054 p	Similar to two-component response regulator
3	3626	1515	SA-2516.2	1615563-1616237 p	similar to putative NAD(P)H-flavin oxidoreductase
36	3627	1516	SA-2517.1	1616325-1616717 p	similar to S-D-lactolyglutathione methylglyoxal lyase
36	3628	1517	SA-2518.1	1616846-1617772 p	similar to putative sugar transferase
8	3629	1519	SA-252.1	309343-310146 p	Similar to amino acid ABC transporter (permease)



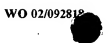
SEQ ID n° 5766	3630	1520	SA-2520.3	152358-154232 p	similar to unknown proteins
SEQ ID n° 5767	3631	1521	SA-2522.2	1486533-1486805 m	30S ribosomal protein S16
SEQ ID n° 5768	3632	1522	SA-2523.2	1486281-1486523 m	similar to unknown protein
SEQ ID n° 5769	3633	1523	SA-2524.2	1484638-1486149 m	similar to transcription regulator RofA related
SEQ ID n° 5770	3634	1524	SA-2525.2	960415-962247 p	similar to GTP binding proteins
SEQ ID n° 5771	3635	1525	SA-2526.2	1364617-1366431 m	Exonuclease motif predicted by PFAM
SEQ ID n° 5772	3636	1527	SA-253.1	308436-309248 p	Similar to amino acids ABC transporter (binding protein)
					similar to a DNA polymerase like protein from Plasmodium
SEQ ID n° 5773	3637	1528	SA-2531.3	1366847-1368082 p	falciparum
SEQ ID n° 5774	3638	1529	SA-2532.1	1062952-1063155 m	Unknown
SEQ ID n° 5775	3639	1530	SA-2533.1	649127-649315 p	Unknown
SEQ ID n° 5776	3640	1531	SA-2534.1	649051-649233 p	bseudogene
SEQ ID n° 5777	3641	1532	SA-2535.1	579004-579279 p	HU like DNA-binding protein
					similar to geranyltranstransferase (farnesyl diphosphate
SEQ ID n° 5778	3642	1533	SA-2536.1	572721-573593 p	synthase)
SEQ ID n° 5779	3643	1534	SA-2537.2	565465-565695 p	similar to unknown proteins
SEQ ID n° 5780	3644		SA-2541.2	2046191-2046337 m	Unknown
SEQ ID n° 5781	3645	1541	SA-2548.1	1263082-1263228 m	similar to unknown proteins
EQ ID n° 5782	3646	1542	SA-2549.1	1264885-1265175 p	Unknown
EQ ID n° 5783	3647	1543	SA-255.1	306955-308298 p	similar to ATP-dependent RNA helicase
SEQ ID n° 5784	3648	1546	SA-2555.2	507784-508122 p	Unknown
SEQ ID n° 5785	3649	1547	SA-2556.1	1622361-1622597 m	Similar to protein-export protein SecG
SEQ ID n° 5786	3650	1549	SA-2558.1	1631019-1631249 m	Unknown
SEQ ID n° 5787	3651	1550	SA-2559.1	659421-659609 m	similar to transposase
0000	0100	, and a second	0		
SEC 10 n° 5789	3653	1551	SA-2560 1	505397-305657 p	similar to prospro-in-acetylmuramoyi-pentapeptide transferase
SEQ ID n° 5790	3654	1559	SA-257.1	303587-305845 p	similar to penicillin-binding protein 2X
SEQ ID n° 5791	3655	1560	SA-2571.3	984145-984786 p	Similar to ABC transporter (permease)
SEQ ID n° 5792	3656	1561	SA-2572.1	983681-984010 p	similar to unknown proteins
SEQ ID n° 5793	3657	1564	SA-2576.2	1649770-1650459 m	ribosomal protein L1
SEQ ID n° 5794	3658	1566	SA-258.1	303257-303583 p	similar to cell division protein FtsL
SEQ ID n° 5795	3659	1567	SA-2582.2	1123914-1124204 m	similar to unknown protein
SEQ ID n° 5796	3660	1568	SA-2583.3	2056908-2057630 m	similar to unknown proteins
SEQ ID n° 5797	3661	1572	SA-259.1	302295-303242 p	similar to unknown proteins
SEQ ID'n° 5798	3662	1575	SA-2596.1	1720081-1720746 m	similar to amino acid ABC transporter (permease)



					Unknown, similar to amino acid ABC transporter (binding
SEQ ID n° 5799	3663	1576	SA-2597.2	1720770-1721627 m	protein)
SEQ ID n° 5800	3664	1577	SA-2598.3	1077491-1078357 m	similar to phosphate ABC transporter (binding protein)
SEQ ID n° 5801	3665	1578	SA-2599.1	1205070-1206410 m	Similar to drug-export protein
SEQ ID n° 5802	3666	1579	SA-26.1	1051977-1052234 m	Unknown
SEQ ID n° 5803	3667	1580	SA-260.1	300959-302212 p	similar to Gamma-glutamyl phosphate reductase
SEQ ID n° 5804	3668	1589	SA-261.1	300146-300949 p	similar to gamma-glutamyl kinase
0 0 0 0 0	000	1017	7 000		similar to Similar to beta-glucoside specific PTS system enzyme
SEC 10 n 3803	3006	1595	SA-263.1	298620-300005 m	IIBC
SEQ ID n° 5806	3670	1596	SA-2632.1	1064917-1065144 m	Unknown
SEQ ID n° 5807	3671	1598	SA-264.1	296808-298412 p	Transmembrane protein similar to unknown proteins
SEQ ID n° 5808	3672	1602	SA-265.1	296070-296804 p	Similar to ABC transporter (ATP-binding protein)
SEQ ID n° 5809	3673	1604	SA-2651.1	2042706-2042933 p	Hypothetical gene
SEQ ID n° 5810	3674	1606	SA-2655.1	81684-82052 p	ribosomal protein L14
SEQ ID n° 5811	3675	1608	SA-266.1	295745-296050 p	similar to unknown proteins
SEQ ID n° 5812	3676	1613	SA-267.1	293547-295532 p	similar to transketolase
SEQ ID n° 5813	3677	1616	SA-2673.1	827405-827671 m	Similar to unknown proteins
SEQ ID n° 5814	3678	1617	SA-268.1	291992-293422 p	similar to transcriptional regulatory protein (N-terminal part)
SEQ ID n° 5815	3679	1622	SA-269.1	290664-292001 p	similar to NADH oxidase
SEQ ID n° 5816	3680	1624	SA-2691.1	2150868-2151116 p	similar to phage protein
SEQ ID n° 5817	3681	1626	SA-270.1	289878-290576 p	similar to glycerol uptake facilitator
SEQ ID n° 5818	3682	1627	SA-2703.1	472009-472830 p	similar to unknown protein
SEQ ID n° 5819	3683	1629	SA-2707.1	1510882-1511082 m	50S ribosomal protein L35
SEQ ID n° 5820	3684	1630	SA-2709.1	1511122-1511652 m	translation initiation factor IF-3
SEQ ID n° 5821	3685	1631	SA-271.1	288037-289866 p	glycerol-3-phosphate dehydrogenase
SEQ ID n° 5822	3686	1632	SA-2710.1	1511813-1512496 m	Similar to cytidine monophosphate kinase
SEQ ID n° 5823	3687	1634	SA-2712.1	1512507-1513031 m	Similar to unknown proteins (serine rich)
SEQ ID n° 5824	3688	1638	SA-272.1	286516-288024 p	glycerol kinase
SEQ ID n° 5825	3689	1641	SA-273.1	286139-286402 p	similar to unknown proteins
SEQ ID n° 5826	3690	1643	SA-274.1	285793-286050 p	similar to unknown proteins
SEQ ID n° 5827	3691	1646	SA-275.1	283742-285781 p	glycyl-tRNA synthetase (beta subunit)
SEQ ID n° 5828	3692	1648	SA-2756.1	1714969-1715208 m	Similar to unknown proteins
SEQ ID n° 5829	3693	1651	SA-2759.1	1715366-1715908 m	Similar to unknown proteins
SEQ ID n° 5830	3694	1652	SA-276.1	283097-283738 p	similar to acyl carrier protein phosphodiesterase
SEQ ID n° 5831	3695	1655	SA-2763.2	1878992-1880002 m	glyceraldehyde 3-phosphate dehydrogenase
SEQ ID n° 5832	3696	1658	SA-2768.2	1349651-1350061 m	similar to unknown proteins



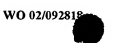
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1666 SA-278.2 281328-281870 p
SA-2787.3 1919967-1920950 m
SA-2789.1 508125-508481 p
SA-2799.2
1668 SA-28.1 730066-732432 p
1669 SA-280.2 1336040-1339798 m
_
SA-2803.2 509314-510102 p
SA-2804.2 1613541-1614263 m
2
SA-2815.2
1670 SA-282.1 1334988-1335905 m
2
SA-2829.1
-
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2
SA-2843.1 716495-717295 m
SA-2849.1
13:
SA-2855.1 1363655-1364491 p



SEQ ID n° 5867	3731		SA-2856.1	1493128-1493649 m	similar to pyrimidine biosynthetic operon repressor
SEQ ID n° 5868	3732		SA-2857.2	1557873-1558250 p	large conductance mechanosensitive channel protein MscL
SEQ 1D n° 5869	3733	1673	SA-286.1	1332440-1333678 m	similar to GTP-binding protein
SEQ ID n° 5870	3734		SA-2860.1	1169149-1169730 m	similar to xanthine phosphoribosyltransferase
SEQ ID n° 5871	3735		SA-2861.1	1245656-1245853 p	similar to hypothetical transcriptional regulator
ို	3736		SA-2862.1	1251589-1251837 m	similar to unknown proteins
SEQ ID n° 5873	3737		SA-2863.1	1251885-1252130 m	similar to unknown proteins
SEQ ID n° 5874	3738		SA-2866.1	762834-763217 p	similar to putative plasmid replication protein
SEQ ID n° 5875	3739		SA-2867.1	762543-762821 p	Unknown
SEQ ID n° 5876	3740		SA-2868.1	761728-762546 p	similar to plasmid partition protein ParA
SEQ ID n° 5877	3741		SA-2869.1	761005-761589 p	Unknown
SEQ ID n° 5878	3742	1674	SA-287.1	1331800-1332447 m	similar to unknwon proteins
SEQ ID n° 5879	3743		SA-2870.1	759383-759640 p	Unknown
SEQ ID n° 5880	3744		SA-2871.1	757414-759363 p	similar to plasmid protein
SEQ ID n° 5881	3745		SA-2872.1	755988-756221 p	Unknown
SEQ ID n° 5882	3746		SA-2874.1	753724-755931 p	similar to unknown protein
SEQ ID n° 5883	3747		SA-2875.1	753122-753604 p	Unknown
SEQ ID n° 5884	3748		SA-2877.1	749873-753022 p	similar to plasmid protein
SEQ ID n° 5885	3749		SA-2878.1	749623-749880 p	Unknown
SEQ ID n° 5886	3750		SA-2879.1	748964-749395 p	similar to single-strand binding protein
SEQ ID n° 5887	3751	1675	SA-288.1	1330829-1331758 m	Similar to unknown proteins
SEQ ID n° 5888	3752		SA-2880.1	1036003-1038804 m	similar to surface antigen proteins, putative peptidoglycan bound protein (LPXTG motif)
					similar to unknown protein, putative peptidoglycan bound protein
SEQ ID n° 5889	3753		SA-2882.1	745217-745939 p	(LPXTG motif)
SEQ ID n° 5890	3754		SA-2883.1	742879-745140 p	Putative peptidoglycan bound protein (LPXTG motif)
SEQ ID n° 5891	3755		SA-2885.1	741782-742864 p	Unknown
SEQ ID n° 5892	3756		SA-2886.1	741424-741771 p	6 No similarity
					similar to ATP-dependent Clp protease ATP-binding subunit
SEQ ID n° 5893	3757		SA-2887.1	739413-741332 p	ClpA
SEQ ID n° 5894	3758		SA-2888.1	737750-739498 p	similar to DNA topolsomerase III
SEQ ID n° 5895	3759		SA-2890.1	406767-411497 p	similar to plasmid proteins
SEQ ID n° 5896	3760		SA-2891.1	732558-732815 p	Unknown
SEQ ID n° 5897	3761		SA-2893.1	1052360-1054726 m	similar to unknown proteins
SEQ ID n° 5898	3762		SA-2894.1	729554-730012 p	Unknown
SEQ ID n° 5899	3763		SA-2895.1	1055255-1056055 m	similar to unknown proteins



SEO ID 5° 6900	3764		CA 2006 4	2 707007 000007	Current I
SEO ID nº 5901	3765		SA-2897 1	728057-08057	Takaowa
SEQ ID n° 5902	3766	1676	SA-29.1	1054780-1055238 m	Unknown
SEQ ID n° 5903	3767	1677	SA-290.1	1330066-1330827 m	similar to oxidoreductase
SEQ ID n° 5904	3768		SA-2900.1	727411-727686 p	Unknown
SEQ ID n° 5905	3769		SA-2902.1	726446-727288 p	hypothetical start codon
SEQ 1D n° 5906	· 3770		SA-2903.1	- 725380-726384 p	Unknown
SEQ ID n° 5907	3771		SA-2905.1	724345-725301 m	Unknown
SEQ ID n° 5908	3772		SA-2906.1	724104-724352 p	Unknown
SEQ ID n° 5909	3773		SA-2907.1	723919-724089 p	Unknown
SEQ ID n° 5910	3774		SA-2908.1	723358-723768 p	Unknown
					similar to similar to single-strand DNA-specific exonuclease
SEQ ID n° 5911	3775	1678	SA-291.1	1327871-1330069 m	RecJ
SEQ ID n° 5912	3776		SA-2910.1	722916-723365 p	Unknown
SEQ ID n° 5913	3777		SA-2911.1	1062066-1062512 m	similar to unknown proteins
SEQ ID n° 5914	3778		SA-2912.1	721852-722280 p	Unknown
SEQ ID n° 5915	3779		SA-2913.1	721637-721840 p	Unknown
SEQ ID n° 5916	3780		SA-2914.1	1063411-1064904 m	similar to plasmid replication protein
SEQ ID n° 5917	3781		SA-2916.1	719648-719875 p	Unknown
SEQ ID n° 5918	3782		SA-2917.1	719536-719655 p	Unknown
SEQ ID n° 5919	3783		SA-2918.1	719438-719530 p	hypothetical gene
SEQ ID n° 5920	3784		SA-2919.1	719103-719432 p	Unknown
SEQ ID n° 5921	3785	1679	SA-292.1	1325155-1327719 m	similar to putative hydrolytic protein
SEQ ID n° 5922	3786		SA-2921.3	1558627-1559130 m	similar to unknown proteins
SEQ ID n° 5923	3787		SA-2922.1	1559213-1560019 m	Similar to amino acid ABC transporter (binding protein)
SEQ ID n° 5924	3788		SA-2926.1	1882439-1882909 m	ribosomal protein S7
SEQ ID n° 5925	3789		SA-2928.1	1882931-1883344 m	ribosomal protein S12
SEQ ID n° 5926	3790	1680	SA-293.1	1324514-1325032 m	similar to adenine phosphoribosyltransferase
SEQ ID n° 5927	3791		SA-2939.1	1982190-1982534 m	similar to hypothetical thioredoxin
SEQ ID n° 5928	3792	1681	SA-294.1	1323716-1324396 m	similar to unknown proteins
SEQ ID n° 5929	3793		SA-2940.1	1981772-1982113 m	similar to unknown proteins
SEQ ID n° 5930	3794		SA-2945.1	1765545-1765901 m	similar to unknown proteins
SEQ ID n° 5931	3795		SA-2946.1	1701220-1701810 m	ATP-dependent CLP protease proteolytic subunit
SEQ ID n° 5932	3796		SA-2947.1	1700875-1701120 m	similar to unknown proteins
SEQ ID n° 5933	3797	1682	SA-295.1	1322929-1323612 m	similar to unknown proteins
SEQ ID n° 5934	3798		SA-2951.2	1558347-1558523 m	ribosomal protein S21



SEQ ID n° 5935	3799		] SA-2954.1	. 1350730-1351386 p	similar to unknown proteins
SEQ ID n° 5936	3800		SA-2955.1	1350071-1350568 m	similar to unknown proteins
SEQ ID n° 5937	3801	1683	SA-296.1	1322151-1322939 m	Similar to unknown proteins
					similar to N-terminal first 100 amino acids of anaerobic
SEQ ID n° 5938	3802		SA-2961.1	1136745-1137083 m	ribonucleotide reductase
SEQ ID n° 5939	3803	1684	SA-297.1	1321039-1322142 m	similar to probable D-amino acid oxidase
SEQ ID n° 5940	3804		SA-2973.2	433313-433570 p	Unknown
SEQ ID n° 5941	3805		SA-2974.2	431344-433293 p	similar to plasmid transfer complex protein
SEQ ID n° 5942	3806		SA-2975.2	427052-427534 p	Unknown
SEQ ID n° 5943	3807		SA-2976.2	423803-426952 p	similar to plasmid proteins
					similar to surface antigen proteins, putative peptidoglycan bound
SEQ ID n° 5944	3808		SA-2977.2	419921-422722 p	protein (LPXTG motif)
SEQ ID n° 5945	3809		SA-2978.1	718598-719092 p	Unknown
SEQ ID n° 5946	3810		SA-2979.1	718373-718561 p	Unknown
°c	3811	1685	SA-298.5	1320126-1320980 m	glucose-1-phosphate thymidyltransferase
SEQ ID n° 5948	3812		, SA-2985.2	437152-437463 p	Unknown
SEQ ID n° 5949.	3813		SA-2986.2	436764-437147 p	similar to replication initiation protein from Staphylococci plasmid
SEQ ID n° 5950	3814		SA-2987.2	436473-436751 p	Unknown
SEQ ID n° 5951	3815		SA-2988.2	435658-436476 p	similar to plasmid partition protein ParA
SEQ ID n° 5952	3816		SA-2989.1	434935-435519 p	Unknown
SEQ ID n° 5953	3817	1686	SA-299.5	1319519-1320112 m	dTDP-4-keto-6-deoxyglucose-3,5-epimerase
SEQ ID n° 5954	3818		SA-2990.1	433584-434921 p	similar to plasmid proteins
SEQ ID n° 5955	3819		SA-2991.1	430687-431334 p	similar to unknown protein
SEQ ID n° 5956	3820		SA-2992.1	430382-430687 p	Unknown
SEQ ID n° 5957	3821		SA-2993.1	429918-430151 p	Unknown
SEQ ID n° 5958	3822		SA-2995.1	427654-429861 p	similar to unknown proteins
SEQ ID n° 5959	3823		SA-2997.1	423553-423810 p	Unknown
SEQ ID n° 5960	3824		SA-2998.1	422894-423325 p	similar to single-strand binding protein
SEQ ID n° 5961	3825	1687	SA-3.1	1025432-1027381 m	similar to plasmid transfer complex protein
SEQ ID n° 5962	3826	1688	SA-30.1	728737-729537 p	similar to membrane nuclease
SEQ ID n° 5963	3827		SA-3000.1	419147-419869 p	Putative peptidoglycan bound protein (LPXTG motif)
NEO 10 nº 6964	9000		SA 3004 4	- 070070 416800	similar to plasmid surface exclusion protein, putative
2000 10 10 10 10 10 10 10 10 10 10 10 10	2020		04-3001.	416009-419010	
3000 ID II 3800	3029		3A-3002.1	415/12-416/94 p	SIMILIAR TO UTINITIONITY DIOCETTIS
SEC 10 n 2966	3830		SA-3003.1	415354-415/01 p	UNKNOWN

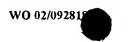


			_		similar to ATP-dependent Clangrage ATP-hinding
SEQ ID n° 5967	3831		SA-3004.1	413430-415262 p	CIPA
SEQ ID n° 5968	3832			411680-413428 p	similar to plasmid DNA topoisomerase
	3833		SA-3007.2	733632-737567 p	similar to plasmid proteins
SEQ ID n° 5970	3834		SA-3008 1	406491-406748 p	Unknown
SEQ ID n° 5971	3835	1689	SA-301.1	1318266-1319312 m	similar to dTDP-glucose-4,6-dehydratase
SEQ ID n° 5972	3836		SA-3010.1	403999-406365 p	similar to unknown proteins
SEQ ID n° 5973	. 3837		SA-3011.1	403487-403945 p	Unknown
SEQ ID n° 5974	3838		SA-3012.1	402670-403470 p	similar to unknown proteins
SEQ ID n° 5975	3839		SA-3013.1	402323-402670 p	Unknown
SEQ ID n° 5976	3840		SA-3014.1	402013-402330 p	Unknown
SEQ ID n° 5977	3841		SA-3016.1	401344-401619 p	Unknown
SEQ ID n° 5978	3842		SA-3018.1	400373-401221 p	Unknown
SEQ ID n° 5979	3843		SA-3019.1	399313-400317 p	similar to unknown protein
SEQ ID n° 5980	3844		SA-3021.1	398278-399234 m	Unknown
SEQ ID n° 5981	3845		SA-3022.1	398037-398285 p	Unknown
SEQ ID n° 5982	3846		SA-3023.1	397879-398022 p	Unknown
SEQ ID n° 5983	3847	_	SA-3025.1	397291-397701 p	Unknown
SEQ ID n° 5984	3848		SA-3027.1	396849-397298 p	Unknown
SEQ ID n° 5985	3849		SA-3028.1	396213-396659 p	similar to unknown proteins
SEQ ID n° 5986	3850		SA-3029.1	395785-396213 p	Unknown
SEQ ID n° 5987	3851		SA-3030.1	395570-395773 p	Unknown
SEQ ID n° 5988	3852		SA-3031.1	393821-395314 p	similar to plasmid replication protein
SEQ ID n° 5989	3853		SA-3033.1	393581-393808 p	Unknown
SEQ ID n° 5990	3854		SA-3034.1	393469-393588 p	Unknown
SEQ ID n° 5991	3855		SA-3035.1	393284-393463 p	Unknown
SEQ ID n° 5992	3856		SA-3036.1	393036-393365 p	Unknown
SEQ ID n° 5993	3857	1691	SA-305.1	1314996-1318214 p	hyaluronate lyase
SEQ ID n° 5994	3858	1692	SA-306.1	1314297-1314779 m	similar to other proteins
SEQ ID n° 5995	3859	1693	SA-307.1	1313846-1314307 m	similar to unknown proteins
SEQ ID n° 5996	3860		SA-3071.1	917842-918033 p	similar to unknown protein
SEQ ID n° 5997	3861		SA-3072.1	733433-733630 p	truncated, C-terminal part
SEQ ID n° 5998	3862		SA-3073.1	732834-733436 p	similar to unknwon proteins (N-terminal part)
SEQ ID n° 5999	3863	1694	SA-308.1	1312564-1313754 m	similar to unknwon proteins
SEQ ID n° 6000	3864	1695	SA-309.1	1311348-1312574 m	similar to unknown proteins
SEQ ID n° 6001	3865	1696	SA-31.1	1056055-1056402 m	Unknown

SEQ ID n° 6002	3866	1697	SA-311.1	1309556-1311238 m	similar to alpha-acetolactate synthase
SEQ ID n° 6003	3867	1698	SA-312.1	1308823-1309542 m	similar to alpha-acetolactate decarboxylase
SEQ ID n° 6004	3868		SA-3125.1	83021-83206 p	ribosomal protein S14
SEQ ID n° 6005	3869		SA-3126.1	88552-88668 p	ribosomal protein L36
SEQ ID n° 6006	3870	1699	SA-313.2	1307114-1308769 p	similar to hypothetical fibronectin-binding protein
SEQ ID n° 6007	3871	1700	SA-314.2	1305752-1306756 m	similar to unknown prorteins
SEQ ID n° 6008	3872	1701	SA-315.1	1304876-1305739 m	Similar to ABC transporter (permease)
SEQ ID n° 6009	3873	1702	SA-316.1	1304115-1304876 m	Similar to ABC transporter (ATP-binding protein)
SEQ 1D n° 6010	3874	1703	SA-318.1	1302157-1303818 m	similar to unknown proteins
SEQ ID n° 6011	3875	1704	SA-319.1	1301312-1302100 m	similar to tributyrin esterase
SEQ ID n° 6012	3876	1705	SA-32.1	1056395-1056712 m	Unknown
SEQ ID n° 6013	3877	1706	SA-320.1	1300512-1301222 m	similar to unknown proteins
SEQ ID n° 6014	3878	1708	SA-322.1	1299603-1300274 m	similar to ribose 5-phosphate isomerase
SEQ ID n° 6015	3879	1709	SA-323.1	1298335-1299546 m	similar to phosphopentomutase
SEQ ID n° 6016	3880	1710	SA-324.1	1297877-1298284 m	similar to arsenate reductase (hypothetical)
SEQ ID n° 6017	3881	1711	SA-325.1	1297029-1297838 m	similar to purine nucleoside phosphorylase
ID n° 6018	3882	1712	SA-326.1	1295771-1297027 m	putative transport protein
SEQ ID n° 6019	3883	1713	SA-327.1	1295077-1295787 m	similar to purine-nucleoside phosphorylase
ID n° 6020	3884	1714	SA-328.1	1294301-1295068 m	similar to unknown proteins
SEQ ID n° 6021	3885	1715	SA-329.2	1293353-1294276 p	similar to transcriptional regulator (LysR family)
SEQ ID n° 6022	3886	1716	SA-330.2	60992-61249 p	similar to unknown proteins
SEQ ID n° 6023	3887	1111	SA-331.2	59909-60982 p	similar to phosphoribosylaminolmidazole carboxylase II
SEQ 1D n° 6024	3888	1718	SA-333.1	59431-59922 p	similar to phosphoribosylaminolmidazole carboxylase
SEQ 1D n° 6025	3889	1719	SA-334.1	57888-59150 p	similar to phosphoribosylamine-glycine ligase
SEQ ID n° 6026	3890	1720	SA-335.1	56801-57604 m	similar to unknown proteins
SEQ ID n° 6027	3891	1721	SA-336.1	55801-56778 p	similar to acetyl xylan esterase (hypothetical)
SEQ ID n° 6028	3892	1722	SA-337.1	54912-55793 p	similar to glucose kinase and to XyIR transcriptional regulator
SEQ ID n° 6029	3893	1723	SA-338.1	53978-54895 p	similar to N-acetylneuraminate lyase
SEQ ID n° 6030	3894	1724	SA-339.1	53319-53981 p	similar to unknown proteins
ID n° 6031	3895	1725	SA-34.1	1057106-1057381 m	Unknown
ID n° 6032	9886	1726	SA-340.1	52856-53299 p	similar to unknown proteins
SEQ ID n° 6033	3897	1727	SA-342.1	52013-52843 p	Similar to ABC transporter (permease)
ID n° 6034	3898	1728	SA-343.1	51116-52003 p	Similar to sugar ABC transporter (permease)
SEQ ID n° 6035	3899	1729	SA-344.1	49712-51028 p	Similar to ABC transporter (binding protein)
ID n° 6036	3900	1730	SA-345.1	48967-49665 p	similar to unknown proteins
	222	22.		7 222 1222 L	



CCO 10 20 CO 27	7000	4704	7 17 7 7 7	- 00701 01171	
SECTION 603/	3901	1731	SA-347.1	4/416-48/20 p	group & streptococcal surrace inmunogenic protein
200	2000	701-	0.010	1 207 1 1 0 1001	similar to bifunctional
					phosphoribosylaminoimidazolecarboxamide
SEQ ID n° 6039	3903	1733	SA-351.1	44630-46177 p	formyltransferase/IMP cyclohydrolase
SEQ ID n° 6040	3904	1734	SA-352.1	43858-44610 p	similar to unknown proteins
SEQ ID n° 6041	3905	1735	SA-353.1	43287-43835 p	similar to Phosphoribosylglycinamide formyltransferase
SEQ ID n° 6042	3906	1736	SA-356.1	42097-43119 p	similar to phosphoribosylformylglycinamide cyclo-ligase
SEQ ID n° 6043	3907	1737	SA-357.1	40615-42069 p	similar to phosphoribosylpyrophosphate amidotransferase
SEQ ID n° 6044	3908	1738	SA-36.1	1057504-1058352 m	Unknown
SEQ ID n° 6045	3909	1740	SA-363.1	36656-40381 p	similar to Phosphoribosylformylglycinamidine synthase
SEO 10 nº 6046	3910	1741	C A 36.4 2	35870,36532	Similar to prosprioribosylaminormoazoresuccinocarboxamide
SEO ID nº 6047	3911	1742	SA-366.2	562006,565167 m	similar to ATD-dependent protesse CinE
SEO ID nº 6048	3012	1743	2000.00	462265 462720 m	Cimilar to undrawan proteins
SEO ID n° 6049	3913	1744	SA-368 1	561899-562201 m	similar to unknown proteins
SEQ ID n° 6050	3914	1745	SA-37.1	1058408-1059412 m	Similar to unknown protein
SEQ ID n° 6051	- 3915	1746	SA-370.1	558990-561782 p	isoleucyl-tRNA synthetase
SEQ ID n° 6052	3916	1747	SA-371.1	557935-558705 p	similar to cell division protein DivIVA
SEQ ID n° 6053	3917	1748	SA-372.1	557137-557925 p	similar to unknown proteins
SEQ ID n° 6054	3918	1749	SA-373.1	556881-557135 p	Similar to unknown proteins
SEQ ID n° 6055	3919	1750	SA-374.1	556273-556878 p	Similar to unknown proteins
SEQ ID n° 6056	3920	1221	SA-375.1	555590-556261 p	Similar to unknown proteins
SEQ ID n° 6057	3921	1752	SA-376.1	554301-555581 p	cell division protein FtsZ
SEQ ID n° 6058	3922	1753	SA-377.1	552906-554279 p	Similar to cell division protein FtsA
SEQ ID n° 6059	3923	1754	SA-379.1	551497-552633 p	Similar to cell division protein DivIB
					Similar to undecaprenyl-PP-MurNAc-pentapeptide-UDPGlcNAc
SEQ ID n° 6060	3924	1755	SA-380.1	550417-551493 p	GlcNAc transferase
SEQ ID n° 6061	3925	1756	SA-381.1	549059-550414 p	Similar to UDP-N-acetylmuramov/alanine0-olutamate linase
SEQ ID n° 6062	3926	1757	SA-382.1	548684-548929 p	Similar to unknown proteins
SEQ ID n° 6063	3927	1758	SA-384.1	546798-548639 p	similar to putative GTP-binding elongation factor
SEQ ID n° 6064	3928	1759	. SA-385.1	546186-546566 p	similar to unknown proteins
SEQ ID n° 6065	3929	1760	SA-387.1	545206-546174 p	similar to glucose kinase
SEQ ID n° 6066	3930	1761	SA-388.1	545000-545209 p	similar to unknown proteins



SEQ ID n° 6067	3931	1762	SA-389.1	544450-544845 m	similar to unknown proteins
SEQ ID n° 6068	3932	1763	SA-39.1	1059491-1060447 p	Unknown
SEQ ID n° 6069	3933	1764	SA-390.1	543817-544449 p	similar to endonuclease III
SEQ ID n° 6070	3934	1765	SA-391.1	542475-543704 p	similar to other proteins
SEQ ID n° 6071	3935	1766	SA-392.1	541439-542482 p	Similar to Acetyl-COA acetyltransferase (truncated)
SEQ ID n° 6072	3936	1767	SA-393.1	540863-541378 p	Unknown
SEQ ID n° 6073	3937	1768	SA-394.1	539746-540738 m	similar to biotin synthetase
SEQ ID n° 6074	3938	1769	SA-395.1	539206-539745 m	similar to other proteins
					similar to Anthranilate synthase component II (Glutamine amido-
SEQ ID n° 6075	3939	1770	SA-396.1	538572-539138 p	(transferase)
SEQ ID n° 6076	3940	1771	SA-397.1	536705-538444 p	isimilar to ABC transporter (exporter) (ATP-binding protein)
Q	3941	1772	SA-398.1	534970-536715 p	similar to ABC transporter (exporter) (ATP-binding protein)
Ω	3942	1773	SA-399.1	534259-534774 p	Similar to unknown proteins
SEQ ID n° 6079	3943	1774	SA-4.2	1027391-1028038 m	similar to unknown protein
SEQ ID n° 6080	3944	1775	SA-40.1	1060440-1060688 m	Unknown
SEQ ID n° 6081	3945	1776	SA-400.1	533153-534259 p	similar to unknown proteins
SEQ ID n° 6082	3946	1777	SA-401.2	532524-533102 p	similar to unknown proteins
SEQ ID n° 6083	3947	1778	SA-402.2	531121-532455 p	Similar to unknown proteins
SEQ ID n° 6084	3948	1779	SA-403.1	529973-531046 p	similar to unknown proteins
SEQ 1D n° 6085	3949	1780	SA-405.1	529498-529983 p	similar to phosphopantetheine adenyltransferase
SEQ ID n° 6086	3950	1781	SA-407.1	529211-529501 p	Unknown
SEQ ID n° 6087	3951	1782	SA-409.1	528447-529199 p	similar to unknown proteins
SEQ ID n° 6088	3952	1783	SA-41.1	1060703-1060846 m	Unknown
SEQ ID n° 6089	3953	1784	SA-410.1	528080-528529 m	Similar to unknown proteins
SEQ ID n° 6090	3954	1785	SA-411.2	527043-528035 p	Similar to asparagine synthetase
SEQ ID n° 6091	3955	1786	SA-412.2	1663493-1664140 p	similar to metal-dependent transcriptional regulator
					Similar to 5 -methylthioadenosine nucleosidase/S-
SEQ ID n° 6092	3956	1787	SA-413.1	1664180-1664869 m	adenosylhomocysteine nucleosidase
SEQ ID n° 6093	3957	1788	SA-414.1	1664879-1665148 m	Similar to unknown proteins
SEQ ID n° 6094	3958	1789	SA-415.1	1665148-1665702 m	Similar to ADP-ribose pyrophosphatase
SEQ ID n° 6095	3959	1790	SA-416.1	1665723-1667102 m	similar to UDP-N-acetylglucosamine pyrophosphorylase
SEQ ID n° 6096	3960	1791	SA-417.1	1667346-1667759 m	Similar to unknown proteins
SEQ ID n° 6097	3961	1792	SA-418.1	1667772-1668149 m	Similar to unknown proteins
SEQ ID n° 6098	3965	1793	SA-419.1	1668238-1669194 m	similar to oxidoreductase
SEQ 10 n° 6099	3963	1794	SA-42.1	1061024-1061434 m	Unknown
SEQ ID n° 6100	3964	1795	SA-420.1	1669191-1669445 m	Isimilar to unknown proteins C-terminal end



SEQ ID n° 6101	3965	1797	SA-422.1	1669406-1669648 m	Unknown
SEQ ID n° 6102	3966	1799	SA-424.1	1669887-1670585 m	similar to oxidoreductases
SEQ ID n° 6103	3967	1800	SA-425.1	1670578-1670814 m	Similar to unknown proteins
SEQ ID n° 6104	3968	1801	SA-426.1	1670904-1671152 m	Similar to unknown proteins
SEQ ID n° 6105	3969	1802	SA-427.1	1671350-1671850 m	similar to other proteins
SEQ ID n° 6106	3970	1803	SA-428.1	1672090-1672545 m	similar to unknown proteins (C-terminal end)
SEQ ID n° 6107	3971	1804	SA-429.1	1672546-1673502 m	Similar to other proteins (N-terminal part)
SEQ 1D n° 6108	3972	1805	SA-430.1	1673642-1675801 m	Similar to unknown proteins
SEQ ID n° 6109	3973	1806	SA-431.1	1675808-1677253 m	Similar to other proteins
SEQ ID n° 6110	3974	1807	SA-432.1	1677246-1677923 m	Unknown
SEQ ID n° 6111	3975	1808	SA-433.1	1677975-1678586 m	Unknown
SEQ ID n° 6112	3976	1809	SA-434.2	1679034-1680371 m	Similar to branched-chain amino acid transporter
SEQ ID n° 6113	3977	1810	SA-435.1	1680567-1682564 m	methionyl-tRNA synthetase
SEQ ID n° 6114	3978	1811	SA-436.1	1682707-1683582 p	Similar to other proteins
SEQ ID n° 6115	3979	1812	SA-437.1	1683740-1684435 m	Similar to unknown proteins
SEQ ID n° 6116	3980	1813	SA-438.1	1684771-1685988 p	Similar to PTS enzyme IIC
SEQ ID n° 6117	3981	1814	SA-439.1	1685972-1686814 p	Similar to unknown proteins
SEQ ID n° 6118	3982	1815	SA-44.1	1061427-1061876 m	Unknown
SEQ ID n° 6119	3983	1816	SA-441.1	1686891-1687718 p	Similar to 3 -exo-deoxyribonuclease
SEQ.ID n° 6120	3984	1817	SA-442.1	1687759-1688115 m	Similar to other proteins
SEQ ID n° 6121	3985	1818	SA-443.1	1688117-1688593 m	similar to O6-alkylguanine-DNA-alkyltransferase
SEQ ID n° 6122	3986	1819	SA-445.1	1688649-1689830 m	similar to phosphoglycerate dehydrogenase
SEQ ID n° 6123	3987	1820	SA-446.1	1689892-1690440 m	similar to unknown proteins
SEQ ID n° 6124	3988	1821	SA-447.1	1690509-1691600 m	similar to phosphoserine aminotransferase
SEQ ID n° 6125	3989	1822	SA-448.1	1691733-1692368 p	Similar to other proteins
SEQ ID n° 6126	3990	1823	SA-45.1	722280-722726 p	Unknown
SEQ ID n° 6127	3991	1824	SA-450.1	1692638-1693501 m	Similar to unknown proteins
SEQ ID n° 6128	3992	1825	SA-451.1	1693507-1693833 m	Similar to unknown proteins
SEQ ID n° 6129	3993	1826	SA-452.1	1693864-1694727 m	similar to DNA polymerase III (delta subunit)
SEQ ID n° 6130	3994	1827	SA-453.1	1694747-1695382 m	similar to thymidylate kinase
SEQ ID n° 6131	3995	1828	SA-454.1	1695471-1696130 m	Similar to acetoin dehydrogenase
SEQ ID n° 6132	3996	1829	SA-455.1	1696149-1696859 m	similar to amino acld ABC transporter (ATP-binding protein)
SEQ ID n° 6133	3997	1830	SA-456.1	1696859-1697623 m	similar to amino acid ABC transporter (ATP-binding protein)
SEQ ID n° 6134	3998	1831	SA-457.1	1697624-1698577 m	similar to ABC transporter (permease)
SEQ ID n° 6135	3999	1832	SA-458.2	1698580-1699449 m	similar to amino acid ABC transporter (permease)
SEQ ID n° 6136	4000	1833	SA-46.1	1062512-1062940 m	Unknown

19	Unknown	389634-390137 p	SA-495.1	1867
303	similar to cell-cycle regulation histidine triad (HIT) protein	388879-389298 m	SA-494.1	1866
!/U:	similar to unknown protein	388586-388882 m	SA-493.1	1865
302	similar to transcription regulator, hypothetical.	387215-388522 p	SA-491.1	1864
1 / <b>LL</b>	similar to unknown protein	386685-387206 p	SA-490.1	1863
CI	Unknown	1065262-1065354 m	SA-49.2	1862
r	similar to unknown protein	386249-386692 p	SA-489.1	1861
	similar to unknown transmembrane protein	384679-386100 p	SA-488.1	1860
	similar to unknown transmembrane protein	383965-384576 p	SA-486.1	1858
	putative transmembrane protein	383291-383875 p	SA-485.1	1857
	similar to unknown protein	382390-383202 m	SA-484.1	1856
	similar to mannose-specific PTS enzyme IID	381077-382087 m	SA-483.1	1855
4	similar to mannose-specific PTS enzyme IIC	380232-381044 m	SA-481.1	1854
	similar to mannose-specific PTS enzyme IID	379306-380217 m	SA-480.1	1853
	similar to unknown protein	378825-379187 m	SA-478.1	1851
	similar to unknown protein	377720-378787 p	SA-477.1	1850
	seryl-tRNA synthetase	376152-377429 m	SA-475.1	1849
97	[Streptococcus mutans] hypothetical	375474-376106 p	SA-474.1	1848
39	similar to sakacin A production response regulator			
	subunit alpha	374231-375004 p	SA-473.1	1847
	similar to acetyl-coenzyme A carboxylase carboxyl transferase			
	subunit beta	373363-374238 p	SA-472.1	1846
	similar to acetyl-coenzyme A carboxylase carboxyl transferase			
	similar to acetyl-CoA carboxylase biotin carboxylase subunit	371984-373354 p	SA-471.1	1845
	similar to beta-hydroxyacyl-ACP dehydratase	371524-371946 p	SA-470.1	1844
	similar to plasmid replication protein E	719888-721381 p	SA-47.1	1843
	similar to biotin carboxyl carrier protein	371027-371527 p	SA-469.1	1842
	similar to 3-oxoacyl-acyl-carrier protein synthase	369793-371025 p	SA-468.1	1841
	similar to beta-ketoacyl-ACP reductase	369043-369777 p	SA-467.1	1840
	similar to malonyl CoA-acyl carrier protein transacylase	368108-369034 p	SA-466.1	1839
201	similar to putative trans-2-enoyl-ACP reductase II	367129-368088 p	SA-465.1	1838
072	similar to acyl carrier protein	366750-366974 p	SA-464.1	1837
U <i>4</i> ,1	similar to beta-ketoacyl-ACP synthase III	365721-366692 p	SA-463.1	1836
0	similar to transcriptional regulator (MarR family)	365287-365721 p	SA-462.1	1835
**	acid-binding protein	1699555-1700721 m	SA-460.3	1834
	similar to branched-chain amino acid ABC transporter, amino			

4004 4005 4006 4007 4008

SEQ ID n° 6143

SEQ ID nº 6144 SEQ ID n° 6145

SEQ ID n° 6139 SEQ ID n° 6140 SEQ ID n° 6141 SEQ ID n° 6142

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SEQ ID n° 6147 SEQ ID n° 6148

SEQ ID n° 6146

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SEQ ID n° 6149

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SEQ ID n° 6150

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SEQ ID n° 6162

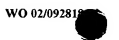
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SEQ ID n° 6168 SEQ ID n° 6167



SEQ ID n° 6169	4033	1868	SA-497.2	1252542-1254821 m	similar to ATP-dependent DNA helicase
SEQ ID n° 6170	4034	1869	SA-498.1	1254927-1255313 m	similar to unknown proteins
SEQ ID n° 6171	4035	1870	SA-499.1	1255446-1256771 m	similar to uracil permease
SEQ ID n° 6172	4036	1871	SA-5.2	1028038-1028343 m	Unknown
SEQ ID n° 6173	4037	1872	SA-50.1	1065137-1065256 m	Unknown
SEQ ID n° 6174	4038	1873	SA-500.1	1257368-1258714 p	similar to probable amino-acid transporter
SEQ ID n° 6175	4039	1874	SA-501.1	1258778-1260013 p	similar to putative cation efflux system protein
SEQ ID n° 6176	4040	1875	SA-502.1	1260172-1260564 p	similar to unknown proteins
SEQ ID n° 6177	4041	1876	SA-503.2	1260545-1261240 p	similar to unknown proteins
SEQ ID n° 6178	4042	1877	SA-504.2	1261309-1261932 p	similar to unknown proteins
SEQ ID n° 6179	4043	1878	SA-505.1	1262334-1262480 m	Unknown
SEQ ID n° 6180	4044	1879	SA-506.1	1262533-1262964 p	lUnknown
SEQ ID n° 6181	4045	1880	SA-507.1	1263261-1263695 p	Unknown
SEQ ID n° 6182	4046	1881	SA-508.2	1264088-1264876 p	similar to repressor protein - phage associated
SEQ ID n° 6183	4047	1882	SA-509.2	1211420-1212304 m	Similar to putative pseudouridine synthase
SEQ ID n° 6184	4048	1883	SA-51.1	1065360-1065689 m	Unknown
SEQ ID n° 6185	4049	1884	SA-510.1	1212301-1213137 m	Similar to unknown protein
SEQ ID n° 6186	4050	1885	SA-511.1	1213112-1213783 m	Similar to unknown protein
SEQ ID n° 6187	4051	1886	SA-512.1	1213893-1214465 p	Similar to unknown protein
SEQ ID n° 6188	4052	1887	SA-513.1	1214642-1215616 p	Similar to Phosphoribosylpyrophosphate synthetase
SEQ ID n° 6189	4053	1890	SA-517.1	362167-363519 m	similar to aspartokinase
SEQ ID n° 6190	4054	1891	SA-518.1	363613-364263 p	similar to unknown protein
SEQ ID n° 6191	4055	1892	SA-519.3	364400-365191 p	similar to enoyl-CoA isomerase
SEQ ID n° 6192	4056	1893	SA-52.1	392531-393025 p	Unknown
SEQ ID n° 6193	4057	1894	SA-520.2	507494-507787 p	Similar to unknown proteins
0EO 10 20 6404	4050	1004	400	000000	similar to alpha protein, putative peptidoglycan linked protein
0E0 10 10 0104	4020	1033	0A-023.4	303440-300020 p	
SEC 10 11 6193	4039	1090	SA-324.1	502012-503205 m	similar to transcriptional regulator (Arac/Ayis ramily)
SEC 10 n 0190	4000	1887	2A-526.1	501293-501817 p	similar to transcriptional regulator tetk-tamily
SEC 10 n 619/	4001	1888	SA-527.3	500298-501161 m	similar to cation efflux system protein
SEQ ID n° 6198	4062	1899	SA-528.3	499830-500213 m	similar to similar to oxydoreductases, N-terminal part
SEQ ID nº 6199	4063	1900	SA-529.1	499359-499829 m	similar to oxydoreductase (C-terminal part)
SEQ ID n° 6200	4064	1901	SA-53.1	392306-392494 p	Unknown
SEQ ID n° 6201	4065	1902	SA-530.1	498794-499249 m	similar to alcohol dehydrogenase (N-terminal part)
SEQ ID n° 6202	4066	1903	SA-531.1	498213-498908 m	similar to alcohol dehydrogenase (C-terminal part)
SEQ ID n° 6203	4067	1904	SA-532.1	497808-498197 m	similar to transcriptional regulator (MerR family)



SEQ ID n° 6204	4068	1905	SA-533.1	497403-497798 m	Similar to other proteins
SEQ ID n° 6205	4069	1906	SA-534.1	497062-497379 m	Similar to decarboxylase
SEQ ID n° 6206	4070	1907	SA-535.1	496691-496975 m	Similar to unknown proteins
SEQ ID n° 6207	4071	1908	SA-536.1	495961-496452 m	Similar to hypothetical transcriptional regulators
SEQ ID n° 6208	4072	1909	SA-537.1	495528-495917 p	similar to unknown proteins
SEO ID nº 6209	4073	1910	S A - 5 3 B 1	702348 405645 P	Similar to surface proteins, putative peptidoglycan bound protein
SEQ ID n° 6210	4074	1911	SA-54.2	391173-392213 p	similar to ABC transporter (permease)
SEQ ID n° 6211	4075	1912	SA-540 1	490117-492285 n	Similar to ribonucleoside-diphosphate reductase 2 aloba subunit
SEQ ID n° 6212	4076	1913	SA-541.1	489702-490115 p	Similar to unknwon proteins
SEQ ID n° 6213	4077	1914	SA-542.2	488691-489701 p	similar to ribonucleoside-diphosphate reductase beta chain
SEQ ID n° 6214	4078	1915	SA-544.1	487230-488138 p	similar to rhamnosyltransferase
					Putative peptidoglycan bound protein (LPXTS motif) similar to
SEQ ID n° 6215	4079	1918	SA-547.2	483319-487020 p	C5A peptidase
SEQ ID n° 6216	4080	1919	SA-548.2	482703-483131 p	similar to unknown proteins
SEQ ID n° 6217	4081	1920	SA-549.1	481905-482669 p	Similar to purine nucleoside phosphorylase
SEQ ID n° 6218	4082	1921	SA-55.2	390446-391171 p	similar to ABC transporter (ATP-binding protein)
					Similar to other proteins including hypothetical
SEQ ID n° 6219	4083	1922	SA-550.1	480283-481638 p	methyltransferases
SEQ ID n° 6220	4084	1923	SA-551.1	479408-480184 m	Similar to unknown proteins
SEQ ID n° 6221	4085	1924	SA-552.1	478791-479324 m	Similar to unknown proteins
SEQ ID n° 6222	4086	1925	SA-554.2	1577255-1578580 m	Similar to unknown proteins
SEQ ID n° 6223	4087	1926	SA-555.1	1578573-1580081 m	Similar to putative glucosyl transferase
SEQ ID n° 6224	4088	1927	SA-557.1	1580095-1582482 m	Similar to preprotein translocase secA
SEQ ID n° 6225	4089	1928	SA-558.1	1582469-1583461 m	Similar to unknown proteins
SEQ 1D n° 6226	4090	1929	SA-559.1	1583458-1585017 m	Similar to unknown protein
SEQ ID n° 6227	4091	1931	SA-560.1	1585024-1586568 m	Similar to unknown protein
SEQ ID n° 6228	4092	1932	SA-561.1	1586568-1587797 m	Similar to preprotein translocase secY
SEQ ID n° 6229	4093	1933	SA-562.1	1587921-1589117 m	Similar to hypothetical glycosyl transferase
SEQ ID n° 6230	4094	1934	SA-563.1	1589178-1590053 m	Similar to putative glycosyltransferase
SEQ ID n° 6231	4095	1935	SA-565.1	1590046-1591230 m	Similar to putative glycosyl transferase
SEQ ID n° 6232	4096	1936	SA-566.1	1591220-1592461 m	Similar to putative glycosyl transferase
SEQ ID n° 6233	4097	1937	SA-567.1	1592458-1593663 m	Similar to putative glycosyl transferase
SEQ ID n° 6234	4098	1938	SA-568.1	1593672-1594679 m	Similar to unknown proteins
SEQ ID n° 6235	4099	1940	SA-57.1	2155624-2156007 m	Unknown



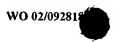
SEQ ID n° 6236	4100	1942	SA-571.1	1595010-1598942 m	Similar to streptococcal hemagglutinin from Streptococcus gordonii, Putative peptidoglycan bound protein (LPXTG motif)
₽	4101	1943	SA-572.1	1599325-1600821 p	Similar to transcription regulator RofA related
°c.	4102	1944	SA-573.1	1600918-1602909 m	excinuclease ABC chain B
SEQ ID n° 6239	4103	1945	SA-574.1	1602970-1603890 m	Similar to unknown proteins
					glutamine ABC transporter permease and substrate binding
SEQ ID n° 6240	4104	1946	SA-575.1	1604073-1606256 p	protein
SEQ ID n° 6241	4105	1947	SA-576.1	1606256-1606996 p	glutamine ABC transporter ATP-binding protein
SEQ ID n° 6242	4106	1948	SA-577.1	1607145-1607495 p	Unknown
SEQ ID n° 6243	4107	1949	SA-578.1	1607587-1607745 m	Similar to unknown proteins
SEQ ID n° 6244	4108	1950	SA-579.1	1607771-1609084 m	Similar to GTP-binding protein
SEQ ID n° 6245	4109	1952	SA-582.1	1609451-1610692 p	Similar to aminopeptidase
SEQ 1D n° 6246	4110	1954	SA-584.1	1610726-1611304 m	Putative peptidoalycan bound serine rich protein (LPXTG motif)
					similar to amidase or hydrolase, putative peptidoglycan bound
SEQ 1D n° 6247	4111	1955	SA-585.4	1611414-1613456 m	protein (LPXTG motif)
SEQ ID n° 6248	4112	1957	SA-589.2	1797911-1798792 m	Similar to fructokinase
SEQ ID n° 6249	4113	1959	SA-590.1	1796846-1797793 m	Similar to mannose-6-phosphate isomerase
SEQ ID n° 6250	4114	1960	SA-591.1	1794209-1796737 m	Similar to preprotein translocase SecA subunit
SEQ ID n° 6251	4115	1961	SA-592.2	1793076-1794083 m	similar to 2-dehydro-3-deoxyphosphoheptonate aldolase
SEQ ID n° 6252	4116	1962	SA-593.2	1792692-1793051 m	similar to holo-acyl-carrier protein synthase
SEQ ID n° 6253	4117	1963	SA-594.2	1791595-1792695 m	similar to alanine racemase
SEQ ID n° 6254	4118	1964	SA-597.1	1789964-1791502 m	Similar to immunogenic secreted protein
SEQ ID n° 6255	4119	1965	SA-598.1	1787869-1789884 m	Similar to ATP-dependent DNA helicase RecG
SEQ ID n° 6256	4120	1966	SA-599.1	1786664-1787578 m	Similar to oxidoreductase
SEQ ID n° 6257	4121	1967	SA-6.1	1028574-1028807 m	Unknown
SEQ ID n° 6258	4122	1968	SA-60.1	2156179-2156709 m	hypothetical gene
SEQ ID n° 6259	4123	1969	SA-600.1	1785688-1786566 m	Similar to shikimate 5-dehydrogenase
SEQ ID n° 6260	4124	1970	SA-601.1	1784691-1785653 p	Similar to L-asparaginase
SEQ ID n° 6261	4125	1971	SA-602.1	1783240-1784622 m	Similar to unknown proteins
SEQ ID n° 6262	4126	1972	SA-603.2	1782732-1783184 p	Similar to unknown proteins
SEQ ID n° 6263	4127	1973	SA-604.2	1781253-1782464 m	Similar to putative aminotransferase
SEQ ID n° 6264	4128	1974	SA-605.1	1780342-1781127 m	Similar to transcriptional regulator (CodY family)
SEQ ID n° 6265	4129	1975	SA-606.1	1779727-1780275 m	Similar to other proteins
SEQ ID n° 6266	4130	1976	SA-607.1	1778715-1779680 p	Similar to 3-hydroxyacyl-CoA dehydrogenase
SEQ ID n° 6267	4131	1977	SA-608.1	1777897-1778526 m	Similar to unknown proteins



SEQ ID n° 6268	4132	1978	SA-609.1	1777056-1777886 m	Similar to unknown proteins
SEQ ID n° 6269	4133	1979	SA-611.1	1774398-1777043 m	Similar to Pyruvate Phosphate Dikinase
SEQ ID n° 6270	4134	1980	SA-612.1	1773958-1774260 m	Similar to Glu-tRNA Gln amidotransferase subunit C
SEQ ID n° 6271	4135	1981	SA-613.1	1772492-1773958 m	Similar to Glutamyl-tRNA Gln amidotransferase subunit A
SEQ ID n° 6272	4136	1982	SA-614.1	1771050-1772492 m	Similar to Glu-tRNA amidotransferase subunit B
SEQ ID n° 6273	4137	1983	SA-615.1	1770006-1770920 m	Similar to unknown proteins
SEQ ID n° 6274	4138	1984	SA-616.1	1769362-1769922 m	Similar to unknown proteins
SEQ ID n° 6275	4139	1985	SA-617.1	1768244-1769362 m	Similar to unknown proteins
ت د	4140	1986	SA-618.1	1767834-1768151 m	Similar to unknown proteins
SEQ ID n° 6277	4141	1987	SA-62.1	2156559-2157413 p	similar to integrase, C-terminal part
SEQ 1D n° 6278	4142	1988	SA-620.1	1767072-1767704 m	Similar to probable nicotinate-nucleotide adenyiyltransferase
SEQ ID n° 6279	4143	1989	SA-621.1	1766488-1767075 m	Similar to unknown proteins
SEQ ID n° 6280	4144	1990	SA-622.2	1765903-1766424 m	similar to unknown proteins
	4145	1991	SA-623.2	818453-819139 p	Similar to unknown proteins
SEQ ID n° 6282	4146	1992	SA-624.1	817104-818324 p	Similar to transporter (antiporter)
					Similar to UDP-N-acetylmuramoylalanyl-D-glutamyl-2,6-
SEQ ID n° 6283	4147	1993	SA-625.1	815550-816917 p	diaminopimelate-D-alanyl-D-alanyl ligase
SEQ ID n° 6284	4148	1994	SA-626.1	814357-815403 p	Similar to D-alanine-D-alanine ligase
SEQ ID n° 6285	4149	1995	SA-627.1	813620-814216 p	Similar to recombination protein RecR
SEQ ID n° 6286	4150	1996	SA-628.1	811563-813605 p	Similar to penicillin binding protein 2B
SEQ ID n° 6287	4151	1997	SA-629.1	810739-811431 p	Similar to phosphoglycerate mutase
SEQ ID n° 6288	4152	1998	SA-63.1	2157770-2158441 p	Unknown
SEQ ID n° 6289	4153	1999	SA-630.1	809804-810562 p	Similar to triosephosphate isomerase
SEQ ID n° 6290	4154	2001	SA-633.1	808427-809623 p	elongation factor Tu
SEQ ID n° 6291	4155	2002	SA-635.1	806807-808075 p	Similar to cell division protein FtsW and to RodA protein
SEQ ID n° 6292	4156	2003	SA-636.1	803906-806701 p	Similar to phosphoenolpyruvate carboxylase
SEQ ID n° 6293	4157	2004	SA-637.1	801898-803697 m	Similar to oligopeptidase
SEQ ID n° 6294	4158	2005	SA-638.1	801450-801839 p	Similar to unknown proteins
SEQ ID n° 6295	4159	2006	SA-639.1	800942-801466 p	Similar to unknown proteins
SEQ ID n° 6296	4160	2007	SA-64.1	2158526-2159197 p	Similar to two-component response regulator
SEQ ID n° 6297	4161	2008	SA-640.1	799937-800785 m	Similar to bacteriophage endolysin
SEQ ID n° 6298	4162	2009	SA-641.1	799287-799904 p	Similar to other proteins
SEQ ID n° 6299	4163	2010	SA-642.1	798519-798992 m	Simllar to transcriptional regulator
SEQ ID n° 6300	4164	2011	SA-643.1	797856-798497 m	Similar to putative phosphoglycerate mutase
SEQ ID n° 6301	4165	2012	SA-644.1	796918-797820 p	Similar to unknown proteins
SEQ 1D n° 6302	4166	2013	SA-646.1	795253-796743 m	lysyl-tRNA synthetase



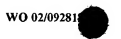
SEQ ID n° 6303	4167	2014	SA-647.1	794708-795178 p	Similar to riboflavin synthase complex beta chain
SEQ ID n° 6304	4168	2015	SA-648.1	793500-794693 p	Similar to GTP cyclohydrase II/ 3,4-dihydroxy-2-butanone-4-phosphate synthase
SEQ ID n° 6305	4169	2016	SA-649.1	792832-793482 p	Similar to ribiflavin synthase alpha chain
SEQ ID n° 6306	4170	2017	SA-65.1	2159181-2160545 p	Similar to two-component sensor histidine kinase
					Similar to riboflavin specific deaminase
					(diaminohydroxyphosphoribosylaminopyrimidine deaminase/5-
SEQ ID n° 6307	4171	2018	SA-651.1	791742-792851 p	amino-6-(5-phosphoribosylamino)uracil reductase)
SEQ ID n° 6308	4172	2019	SA-653.1	789673-791013 m	Similar to manganese transporter
SEQ ID n° 6309	4173	2020	SA-654.2	788774-789571 p	Similar to unknwon proteins
SEQ ID n° 6310	4174	2021	SA-655.2	788438-788650 p	Similar to unknown proteins
SEQ ID n° 6311	4175	2022	SA-656.2	787025-788311 p	Similar to putative peptidases
SEQ ID n° 6312	4176	2023	SA-657.2	d 568987-696387	Similar to putative proteases
SEQ ID n° 6313	4177	2024	SA-658.2	2026535-2028718 m	Similar to glucose-specific PTS enzyme IIABC
SEQ ID n° 6314	4178	2027	SA-660.1	2025666-2026481 m	Similar to unknown protein
SEQ ID n° 6315	4179	2028	SA-661.1	2024637-2025389 m	similar to two-component response regulator
0		0			Similar to two-component sensor histidine kinase (C-terminal
SEQ ID n° 6316	4180	2029	SA-662.1	2024269-2024538 m	part)
SEQ ID n° 6317	4181	2030	SA-663.1	2023298-2024146 m	similar to ABC transporter (ATP-binding protein)
SEQ ID n° 6318	4182	2031	SA-664.1	2022138-2023292 m	Putative transmembrane protein
SEQ ID n° 6319	4183	2033	SA-666.1	2020875-2021927 m	Similar to other proteins, putative transmembrane protein
SEQ ID n° 6320	4184	2034	SA-667.1	2020203-2020628 m	Similar to mannose-specific PTS enzyme IIA
SEQ ID n° 6321	4185	2036	SA-669.1	2019694-2020185 m	Similar to mannose-specific PTS enzyme IIB
SEQ ID n° 6322	4186	2037	SA-67.1	2160655-2162208 m	similar to putative membrane arginine transporter
SEQ ID n° 6323	4187	2038	SA-670.1	2018869-2019678 m	Similar to mannose-specific PTS enzyme IIC
SEQ ID n° 6324	4188	2039	SA-671.1	2018045-2018872 m	Similar to mannose-specific PTS enzyme IID
SEQ ID n° 6325	4189	2040	SA-672.1	2016258-2017907 m	similar to two-component sensor histidine kinase
SEQ ID n° 6326	4190	2041	SA-673.1	2015481-2016254 m	similar to two-component response regulator
SEQ ID n° 6327	4191	2042	SA-674.1	2014432-2015469 m	Similar to iron ABC transporter (binding protein)
SEQ ID n° 6328	4192	2043	SA-675.1	2013713-2014210 p	Similar to unknown proteins
SEQ ID n° 6329	4193	2044	SA-676.1	2012676-2013713 p	Similar to glutamyl-aminopeptidase (hypothetical)
SEQ ID n° 6330	4194	2045	SA-677.1	2012210-2012665 p	Similar to unknown proteins (Nrdt)
SEQ ID n° 6331	4195	2046	SA-678.1	2009651-2012053 p	Similar to nucleotidase (esterase), putative peptidoglycan bound protein (LPXTG motif)
SEQ ID n° 6332	4196	2047	SA-68.1	2162302-2163228 m	similar to carbamate kinase



SEQ ID n° 6333	4197	2048	SA-680.1	2007175-2009391 m	Similar to GTP pyrophosphokinase (stringent response protein ReIA)
SEQ ID n° 6334	4198	2049	SA-681.1	2006722-2007165 m	Similar to unknown proteins
SEQ ID n° 6335	4199	2050	SA-682.1	2005544-2006467 m	Similar to adhesion proteins
SEQ ID n° 6336	4200	2051	SA-686.1	2004072-2005517 m	similar to pneumococcal histidine triad protein B precursor (N-terminal part)
SEQ ID n° 6337	4201	2052	SA-687.1	2002936-2004174 m	similar to pneumococcal histidine triad protein B precursor (C-
SEQ ID n° 6338	4202	2053	SA-688.2	2001878-2002654 p	Similar to transcriptional regulator, DeoR family
SEQ ID n° 6339	4203	2054	SA-689.2	712522-713226 p	similar to two-component response regulator
SEQ ID n° 6340	4204	2055	SA-69.1	2163240-2164238 m	similar to ornithine carbamoyltransferase
SEQ ID n° 6341	4205	2056	SA-690.1	710357-712300 p	theronyl-tRNA synthetase
SEQ ID n° 6342	4206	2057	SA-691.1	708566-709900 p	similar to glucosyl transferase
SEQ ID n° 6343	4207	2058	SA-692.1	707566-708564 p	similar to hexosyltransferase
SEQ ID n° 6344	4208	2059	SA-693.1	706055-707521 p	similar to alpha-amylase
SEQ ID n° 6345	4209	2060	SA-694.2	704919-705923 p	catabolite control protein A
SEQ ID n° 6346	4210	2061	SA-696.2	703624-704709 m	similar to X-Pro dipeptidase
SEQ ID n° 6347	4211	2062	SA-697.1	701776-703566 p	similar to beta-N-acetylglucosaminidase
SEQ ID n° 6348	4212	2063	SA-698.1	700948-701760 p	Unknown
SEQ ID n° 6349	. 4213	2064	SA-699.1	699956-700795 p	similar to oxidoreductase
SEQ ID n° 6350	4214	2066	SA-70.1	2164341-2165636 p	Similar to hypothetical two-component sensor histidine kinase
SEQ ID n° 6351	4215	2067	SA-700.1	698786-699832 p	similar to D-mannonate hydrolase
SEQ ID n° 6352	4216	2068	SA-701.1	697368-698768 p	similar to glucuronate isomerase
SEQ ID n° 6353	4217	2069	SA-702.1	696734-697351 p	similar to 2-dehydro-3-deoxyphosphogluconate aldolase
SEQ ID n° 6354	4218	2070	SA-704.1	695946-696617 p	similar to transcriptional regulator (GntR family)
SEQ ID n° 6355	4219	2071	SA-705.1	694118-695917 p	similar to beta-glucuronidase
SEQ ID n° 6356	4220	2072	SA-706.1	693076-694101 p	similar to 2-keto-3-deoxygluconate kinase
SEQ ID n° 6357	4221	2073	SA-707.1	691459-693009 p	similar to transporter
SEQ ID n° 6358	4222	2074	SA-708.2	690261-691253 p	similar to D-lactate dehydrogenase
SEQ ID n° 6359	4223	2075	SA-71.1	2165633-2166466 p	Similar to hypothetical two-component response regulator
SEQ ID n° 6360	4224	2076	SA-710.2	689189-690247 p	similar to PTS enzyme IIBC
SEQ ID n° 6361	4225	2077	SA-712.1	688112-688996 m	similar to transcriptional regulator, LysR family
SEQ ID n° 6362	4226	2078	SA-713.1	687257-688075 p	Unknown
SEQ ID n° 6363	4227	2079	SA-714.1	686320-687093 p	putative transmembrane protein
SEQ ID n° 6364	4228	2080	SA-715.1	685667-686323 p	similar to ABC transporter (ATP-binding protein)

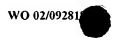


SEQ ID n° 6365	4229	2081	SA-716.1	684812-685450 m	similar to unknown proteins
SEQ ID n° 6366	4230	2082	SA-717.2	683949-684734 p	similar to competence associated membrane nuclease
SEQ ID n° 6367	4231	2083	SA-719.2	1023804-1025141 m	Similar to plasmid proteins
SEQ ID n° 6368	4232	2084	SA-72.1	2166688-2167410 p	Similar to osmoprotectant ABC transporter (ATP-binding protein)
SEQ ID n° 6369	4233	2085	SA-720.1	1023206-1023790 m	Unknown
SEQ ID n° 6370	4234	2086	SA-721.2	1022249-1023067 m	Similar to plasmid partition protein ParA
SEQ ID n° 6371	4235	2087	SA-722.2	1021974-1022252 m	Unknown
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SEQ ID n° 6372	4236	2088	SA-723.2	1021578-1021961 m	similar to replication initiation protein from Staphylococci plasmid
SEQ ID n° 6373	4237	2089	SA-725.2	1021262-1021573 m	Unknown
SEQ ID n° 6374	4238	2090	SA-726.2	1019800-1021128 m	Unknown
SEQ ID n° 6375	4239	2091	SA-727.1	1018711-1019397 m	similar to unknown protein
SEQ ID n° 6376	4240	2092	SA-728.1	1017948-1018721 m	similar to unknown protein
SEQ ID n° 6377	4241	2093	SA-729.1	1016320-1017915 p	similar to oligopeptide and pheromone binding protein
SEQ ID n° 6378	4242	2096	SA-731.2	1014797-1015867 m	similar to integrase/recombinase
SEQ ID n° 6379	4243	2097	SA-732.1	1013816-1014754 p	similar to two-component sensor histidine kinase
SEQ ID n° 6380	4244	2098	SA-733.1	1013151-1013819 p	similar to two-component response regulator
SEQ ID n° 6381	4245	5099	SA-734.1	1011086-1013041 p	1.2 Transport/binding proteins and lipoproteins
SEQ ID n° 6382	4246	2100	SA-735.1	1010332-1011084 p	similar to ABC transporter (ATP-binding protein)
					similar to Lactococcus lactis nisin-resistance protein C-terminal
SEQ ID n° 6383	4247	2101	SA-736.1	1009866-1010306 p	part
					similar to Lactococcus lactis nisin-resistance protein N-terminal
SEQ ID n° 6384	4248	2102	SA-737.1	1009345-1009884 p	part
SEQ ID n° 6385	4249	2103	SA-738.1	1008691-1009179 m	similar to unknown protein
SEQ ID n° 6386	4250	2104	SA-739.1	1007811-1008659 p	similar to other lipoprotein
					similar to osmoprotectant ABC transporter permease and
SEQ ID n° 6387	4251	2105	SA-74.1	2167413-2168927 p	substrate binding protein
SEQ ID n° 6388	4252	2106	SA-740.1	1006935-1007678 p	similar to unknown protein
SEQ ID n° 6389	4253	2107	SA-741.1	1005523-1006857 p	similar to glucose-inhibited division protein
SEQ ID n° 6390	4254	2108	SA-742.1	1004711-1005409 p	similar to transcriptional regulator (GntR family)
SEQ ID n° 6391	4255	2109	SA-743.2	1002936-1004498 m	similar to GMP synthetase
SEQ ID n° 6392	4256	2110	SA-744.3	439342-440142 p	similar to unknown proteins
SEQ ID n° 6393	4257	2111	SA-745.1	440132-440767 p	similar to unknown protein
SEQ ID n° 6394	4258	2112	SA-746.1	441254-441733 p	similar to unknown proteins





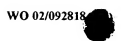
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SEQ ID n° 6427	4291	2150	SA-787.1	638356-639420 m	Similar to unknown proteins
SEQ ID n° 6428	4292	2151	SA-788.1	639649-640932 p	similar to 5-enolpyruvylshikimate-3-phosphate synthase
SEQ ID n° 6429	4293	2152	SA-789.1	640925-641437 p	similar to shikimate kinase
SEQ ID n° 6430	4294	2154	SA-790.1	641494-642867 p	Similar to membrane bound transcriptional regulator
SEQ ID n° 6431	4295	2155	SA-792.1	642968-644323 p	similar to putative RNA methyltransferase
SEQ ID n° 6432	4296	2156	SA-793.1	644431-644652 p	hypothetical CDS
SEQ ID n° 6433	4297	2157	SA-794.1	644770-645507 p	similar to diadenosine tetraphosphatase, acid phosphatase
SEQ ID n° 6434	4298	2158	SA-795.1	645828-646346 p	similar to unknown proteins
					similar to putative transcriptional regulator (TetR/AcrR family) C-
SEQ ID n° 6435	4299	2159	SA-796.1	646475-646711 m	Iterminal part
					similar to putative transcriptional regulator (TetR/AcrR family) N-
SEQ ID n° 6436	4300	2160	SA-797.1	646692-647000 m	terminal part
					similar to C protein alpha-antigen from Streptococcus
SEQ ID n° 6437	4301	2161	SA-798.1	647183-647515 p	agalactiae] N-terminal part
SEQ ID n° 6438	4302	2162	SA-799.1	647636-648562 m	similar to transposase, truncated.
SEQ ID n° 6439	4303	2163	SA-8.1	1028864-1031071 m	similar to unknown proteins
SEQ ID n° 6440	4304	2164	SA-80.1	2175288-2175827 p	similar to transcriptional regulator (TetR/AcrR family)
SEQ ID n° 6441	4305	2165	SA-800.1	648505-648780 m	similar to transposase N-terminal part
SEQ ID n° 6442	4306	2166	SA-801.1	648913-649050 p	similar to unknown protein.
SEQ ID n° 6443	4307	2167	SA-803.1	649438-649785 m	similar to chaperonin (heat shock protein 33 homolog)
SEQ ID n° 6444	4308	2168	SA-804.1	649979-650398 m	similar to transcriptional regulator (C-terminal part)
SEQ ID n° 6445	4309	2169	SA-805.1	650399-651187 m	similar to transcriptional regulator (N-terminal part)
SEQ ID n° 6446	4310	2170	SA-806.1	651570-653234 p	Putative peptidoglycan bound protein (LPXTG motif)
SEQ ID n° 6447	4311	2171	SA-807.1	653323-654246 p	Putative peptidoglycan bound protein (LPXTG motif)
SEQ ID n° 6448	4312	2172	SA-808.1	654248-655165 p	similar to sortase protein
SEQ ID n° 6449	4313	2173	SA-809.3	655122-655973 p	similar to sortase protein
SEQ ID n° 6450	. 4314	2174	SA-81.1	2175913-2176209 m	similar to unknown proteins
SEQ ID n° 6451	4315	2176	SA-811.2	2144199-2144348 p	50S Ribosomal protein L33
SEQ ID n° 6452	4316	2177	SA-812.1	2144001-2144183 p	50S ribosomal protein L32
SEQ ID n° 6453	4317	2178	SA-814.1	2142501-2143781 m	histidyl-tRNA synthetase
SEQ ID n° 6454	4318	2179	SA-817.2	2140657-2142408 m	aspartyl-tRNA synthetase
SEQ ID n° 6455	4319	2180	SA-819.2	2139723-2140667 m	similar to unknown proteins
SEQ ID n° 6456	4320	2181	SA-82.1	2176453-2177064 m	30S ribosomal protein S4
SEQ ID n° 6457	4321	2182	SA-820.1	2138743-2139615 m	similar to unknown proteins
SEQ ID n° 6458	4322	2183	SA-821.1	2138408-2138716 p	similar to unknown proteins
SEQ ID n° 6459	4323	2184	SA-822.1	2136629-2138320 p	arginyl-tRNA synthetase



4325	2186	SA-825.1 SA-826.1	2133337-2135913 m 2133077-2133280 p	Similar to argume repressor Argic DNA mismatch repair protein MutS similar to cold shock protein E
4327	2188	SA-827.1 SA-828.1	2130878-2132851 m 2129614-2130846 m	Similar to DNA mismatch repair Mutt.
4329	2190	SA-829.1	2129022-2129612 m	
4330	2191	SA-83.1	2177394-2177681 m	similar to unknown proteins
4331	2192	SA-831.1	2128448-2128999 m	similar to 3-methyl-adenine DNA glycosylase I
4332	2193	SA-832.1	2127100-2128359 m	similar to competence-damage inducible protein CinA
4333	2194	SA-833.1	2125887-2127026 m	recombination protein RecA
4334	2195	SA-834.1	2125273-2125671 m	similar to unknwon proteins
4335	2196	SA-835.1	2124805-2125071 m	similar to unknwon proteins
4336	2197	SA-837.1	2124386-2124805 m	similar to unknown proteins
4337	2198	SA-838.1	2124043-2124360 m	similar to unknown proteins
4338	2199	SA-839.2	2122252-2123793 m	similar to unknown proteins
4339	2200	SA-84.1	2177693-2179048 m	replicative DNA helicase DnaC
4340	2201	SA-842.3	216500-218530 p	Similar to trehalose-specific PTS enzyme IIABC
4341	2202	SA-843.1	218752-220377 p	similar to trehalose-6-phosphate hydrolase
4342	2203	SA-844.1	220597-222633 p	similar to hypothetical transcriptional antiterminator (BgIG family)
4343	2204	SA-845.1	222636-222920 p	similar to unknown proteins
4344	2205	SA-846.1	222933-224288 p	putative transmembrane protein similar to unknown proteins
4345	2207	SA-848.1	224291-225148 p	similar to other proteins (including putative transketolase)
4346	2208	SA-849.1	225145-226074 p	similar to other proteins (including putative transketolase)
4347	2209	SA-85.1	2179091-2179543 m	50S ribosomal protein L9
4348	2210	SA-850.1	226102-227442 p	similar to unknown proteins
4349	2211	SA-851.1	227530-227799 p	ribosomal protein S15
4350	2212	SA-852.1	228180-230309 p	polynucleotide phosphorylase, alpha chain
4351	2213	SA-853.1	230311-231063 p	similar to unknown protein
4352	2214	SA-854.1	231072-231656 p	similar to serine acetyltransferase
4353	2215	SA-855.1	231666-231848 p	Unknown
4354	2216	SA-856.1	231845-233188 p	cysteinyl-tRNA synthetase
4355	2217	SA-857.1	233181-233567 p	similar to unknown proteins
4356	2218	SA-858.1	233676-234425 p	similar to tRNA/rRNA methyltransferase
4357	2219	SA-859.1	234422-234940 p	similar to unknown protein
4358	0000	CA BG 1	2470EA0 2484E24 m	cimilar to uncodern or cimina



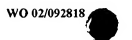
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-87.1 2181598-2183499 m SA-87.1 2181598-2183499 m SA-87.1 193085-193450 p SA-87.1 195623-19584 p SA-87.1 195632-196545 p SA-87.1 195628-195857 p SA-87.1 195632-196545 p SA-88.1 196470-196531 p SA-88.1 196470-196531 p SA-88.1 196470-196531 p SA-88.1 196470-19654 p SA-88.1 196401-199594 p SA-89.1 579717-580649 m SA-89.1 580336-58301 m SA-89.1 580336-583301 m SA-89.1 580336-583301 m SA-89.1 58039-583301 p SA-89.1 580268-591128 m SA-89.1 580268-591128 m SA-89.1 590268-591128 m SA-89.1 590268-591128 m SA-89.1 1031191-1031673 m	similar to tRNA (5-methylaminomethyl-2-thiouridylate)- methyltransferase	2184312-2185433 m	SA-90.1	2255	4392
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-871.1 2181598-2183499 m SA-872.1 189321-192971 p SA-877.1 193085-193450 p SA-877.1 193085-193450 p SA-877.1 193085-193450 p SA-877.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-881.1 196470-196931 p SA-882.1 196470-196931 p SA-882.1 196470-196931 p SA-883.1 196401-199594 p SA-883.1 198401-199594 p SA-884.1 199745-19951 p SA-889.1 579777-580649 m SA-891.1 580836-582071 m SA-891.1 582090-583301 m SA-892.1 584534 m SA-892.1 584534 m SA-893.1 584534-586336 m SA-896.1 586509-583301 m SA-896.1 590268-591128 m SA-899.1 590301-594625 p	Unknown	1031191-1031673 m	SA-9.1	2254	22
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-871 2181598-2183499 m SA-871 193085-193450 p SA-877.1 193085-193450 p SA-877.1 193085-193450 p SA-878.1 193623-196245 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195895 p SA-881.1 196470-196931 p SA-882.1 196470-196931 p SA-883.1 196401-19959 p SA-884.1 199745-199551 p SA-884.1 199745-199951 p SA-889.1 579717-580649 m SA-891.1 582090-583301 m SA-891.1 582090-583301 m SA-892.1 584534 m SA-893.1 584534 m SA-893.1 584534-586733 m SA-895.1 586809-583301 p SA-895.1 586509-583301 m SA-896.1 586509-583301 m	Similar to other proteins	593301-594425 p	SA-899.1	23	2253
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-871.1 2181598-2183499 m SA-872.1 189321-192971 p SA-877.1 193085-193450 p SA-877.1 193085-193450 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195857 p SA-878.1 195628-195895 p SA-881.1 196470-196931 p SA-882.1 196470-196931 p SA-883.1 196401-199594 p SA-883.1 198401-199549 m SA-883.1 199745-199951 p SA-881.1 2183669-2184280 m SA-891.1 582090-583301 m SA-892.1 584534 m SA-892.1 584534 m SA-893.1 584534-586733 m SA-895.1 586509-583301 m	similar to fructose-1,6-bisphosphatase	591280-593211 p	SA-898.1	2252	22
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-871.1 2181598-2183499 m SA-872.1 189321-192971 p SA-877.1 193085-193450 p SA-877.1 193623-196245 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p SA-877.1 195628-19584 p SA-879.1 196217-196516 p SA-881.1 196217-196516 p SA-881.1 196470-196931 p SA-881.1 198401-199594 p SA-882.1 198401-199594 p SA-881.1 199745-19951 p SA-882.1 199745-19951 p SA-881.1 199745-19951 p SA-882.1 579717-580649 m SA-891.1 582090-583301 m SA-891.1 582090-583301 m SA-892.1 58233301 m SA-893.1 584534 m SA-893.1 584534 585346 m SA-895.1 586809-587195 p	similar to unknown proteins	590268-591128 m	SA-897.1	2251	22
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SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-872.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193085-193450 p SA-877.1 195528-195857 p SA-876.1 194440-195531 p SA-876.1 195628-195857 p SA-876.1 196217-196516 p SA-880.1 196470-196931 p SA-881.1 196470-196931 p SA-882.1 196401-199594 p SA-882.1 198401-199594 p SA-885.2 200188-200643 p SA-885.2 200188-200643 p SA-885.2 200188-200643 p SA-885.2 200188-200643 p SA-885.1 2183699-2184280 m SA-889.1 2183669-2184280 m SA-890.1 580836-582071 m	similar to cell wall muropeptide branching enzyme	583314-584534 m	SA-892.1	စ္	224
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193085-193450 p SA-877.1 193623-194594 p SA-877.1 195528-195857 p SA-878.1 195628-195857 p SA-878.1 196217-196516 p SA-881.1 196322-196369 p SA-882.1 196309-197280 p SA-882.1 196470-199594 p SA-884.1 199745-199951 p SA-884.1 199745-199951 p SA-885.2 200188-200643 p SA-886.1 579777-580649 m SA-889.1 2183669-2184280 m SA-89.1 2183669-2184280 m	similar to cell wall muropeptide branching enzyme	582090-583301 m	SA-891.1	5	224
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-877.1 193623-194594 p SA-877.1 195228-195857 p SA-877.1 195228-195857 p SA-878.1 196217-196516 p SA-878.1 196209-197280 p SA-881.1 196909-197280 p SA-881.1 196909-197280 p SA-883.1 196401-199594 p SA-883.1 196401-199594 p SA-883.1 198401-199594 p SA-883.1 579717-580649 m SA-889.1 218369-2184280 m	similar to Cell Wall Muropeptide Branching Enzyme	580836-582071 m	SA-890.1	4	2244
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-877.1 193623-194594 p SA-877.1 195628-195837 p SA-877.1 195628-195837 p SA-877.1 195621-196931 p SA-877.1 196908-197280 p SA-881.1 196908-197280 p SA-882.1 196908-197280 p SA-883.1 196908-19693 p SA-885.2 200188-200643 p SA-887.2 579476-579673 p	similar to unknown proteins	2183669-2184280 m	SA-89.1	_	2243
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193623-194594 p SA-877.1 193623-194594 p SA-877.1 195528-195831 p SA-877.1 195528-195831 p SA-880.1 196470-196931 p SA-881.1 196909-197280 p SA-882.1 196909-197280 p SA-882.1 196401-199594 p SA-883.1 198401-199594 p SA-884.1 196745-199951 p SA-884.1 199745-199951 p SA-885.2 200188-200643 p	similar to dihydroorotate dehydrogenase A	579717-580649 m	SA-889.1		2242
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-877.1 2181598-2183499 m SA-877.1 189321-192971 p SA-877.1 193623-194594 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p SA-878.1 195628-195845 p SA-8879.1 195628-195845 p SA-881.1 196470-196931 p SA-881.1 196470-196931 p SA-881.1 198401-199594 p SA-883.1 198401-199594 p SA-884.1 199745-199591 p	Unknown	579476-579673 p	SA-887.2		2241
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-872.1 189321-192971 p SA-875.1 193085-193450 p SA-875.1 193623-194594 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p SA-877.1 195628-19631 p SA-877.1 195632-196245 p SA-878.1 196470-196931 p SA-880.1 196470-196931 p SA-881.1 198909-19599 p SA-883.1 198401-199594 p	similar to unknown proteins	200188-200643 p	SA-885.2	)	2240
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-872.1 189321-192971 p SA-875.1 193085-194594 p SA-875.1 193623-194594 p SA-877.1 195628-195857 p SA-877.1 195628-196857 p SA-877.1 195632-196545 p SA-878.1 196832-196545 p SA-879.1 196832-196545 p SA-880.1 196909-197280 p SA-882.1 195395-198369 p	similar to transcriptional regulator	199745-199951 p	SA-884.1	9	5239
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193623-194594 p SA-876.1 193623-194594 p SA-876.1 195628-195857 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p SA-878.1 196832-196516 p SA-880.1 196909-197280 p SA-881.1 196909-197280 p	acetate kinase	198401-199594 p	SA-883.1	8	2238
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-866.2 182808-185105 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193623-194594 p SA-875.1 193623-194594 p SA-876.1 194440-195531 p SA-877.1 195528-195857 p SA-877.1 195832-196245 p SA-879.1 196932-196931 p SA-880.1 196909-197280 p	similar to unknown proteins	197395-198369 p	SA-882.1	7	2237
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-87.1 2181598-2183499 m SA-87.1 193025-193450 p SA-875.1 193623-194594 p SA-876.1 194440-195531 p SA-877.1 195528-195857 p SA-877.1 195528-195857 p SA-879.1 195528-196245 p SA-879.1 196217-196516 p	similar to unknown proteins	196909-197280 p	SA-881.1		2236
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-875.1 193085-193450 p SA-875.1 193623-194594 p SA-876.1 195528-195837 p SA-877.1 195528-195857 p SA-878.1 1956217-196516 p	similar to hypothetical competence proteins	196470-196931 p	SA-880.1	5	2235
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-872.1 193085-193450 p SA-875.1 193623-194594 p SA-876.1 194440-195531 p SA-877.1 195628-195857 p SA-877.1 195628-195857 p	similar to unknown proteins	196217-196516 p	SA-879.1		2234
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 193021-192971 p SA-874.1 193085-193450 p SA-876.1 193623-194594 p SA-876.1 193623-194594 p	similar to hypothetical competence proteins	195832-196245 p	SA-878.1		2233
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182608-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-874.1 193085-193450 p SA-876.1 193623-194594 p	similar to exogenous DNA-binding protein comGC	195528-195857 p	SA-877.1		2232
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p SA-874.1 193085-193450 p	probably part of the DNA transport machinery, ComGB protein	194440-195531 p	SA-876.1		2231
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-183204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p	similar to transporter (competence protein)	193623-194594 p	SA-875.1		2230
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m SA-872.1 189321-192971 p	similar to unknown proteins	193085-193450 p	SA-874.1		2229
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p SA-87.1 2181598-2183499 m	RNA polymerase beta -subunit	189321-192971 p	SA-872.1		2228
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p SA-868.1 185629-189204 p	Similar to GidA protein	2181598-2183499 m	SA-87.1		2227
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m SA-866.2 182808-185105 p	RNA polymerase beta-subunit	185629-189204 p	SA-868.1	,	2226
SA-861.1 235033-235893 p SA-863.3 236878-238059 m SA-864.3 181438-182697 m	similar to penicillin-binding protein 1b	182808-185105 p	SA-866.2	4	222
SA-861.1 235033-235893 p SA-863.3 236878-238059 m	tyrosyl-tRNA synthetase	181438-182697 m	SA-864.3	3	222
SA-861.1 235033-235893 p	similar to transposase	236878-238059 m	SA-863.3	7	2222
	Similar to unknown protein	235033-235893 p	SA-861.1		2221



SEQ ID n° 6529	4393	2256	SA-901.1	594579-595592 p	Similar to peptide chain release factor RF-2
SEQ ID n° 6530	4394	2257	SA-902.1	595611-596303 p	Similar to cell division ATP-binding protein FtsE
SEQ ID n° 6531	4395	2258	SA-904.1	596287-597216 p	Similar ic cell-division protein FtsX
SEQ ID n° 6532	4396	2259	SA-905.1	597269-597979 m	Similar to unknown proteins
SEQ ID n° 6533	4397	2260	SA-906.2	597976-598611 m	similar to unknown proteins
SEQ ID n° 6534	4398	2261	SA-907.2	909887-910087 p	H+-transporting ATP synthase c chain
SEQ ID n° 6535	4399	2262	SA-908.2	910120-910836 p	H+-transporting ATP synthase a chain
SEQ ID n° 6536	4400	2263	SA-909.1	910854-911351 p	H+-transporting ATP synthase b chain
SEQ ID n° 6537	4401	2264	SA-91.1	, 2185679-2186347 p	similar to L-serine dehydratase beta subunit
SEQ ID n° 6538	4402	2265	SA-910.1	911351-911887 p	H+-transporting ATP synthase delta chain
SEQ ID n° 6539	4403	2266	SA-911.1	911903-913408 p	H+-transporting ATP synthase alpha chain
SEQ ID n° 6540	4404	2267	SA-912.1	913424-914305 p	H+-transporting ATP synthase gamma chain
SEQ ID n° 6541	4405	2268	SA-913.1	914379-915785 p	H+-transporting ATP synthase beta chain
SEQ ID n° 6542	4406	2269	SA-914.1	915798-916211 p	H+-transporting ATP synthase epsilon chain
SEQ ID n° 6543	4407	2270	SA-916.1	916569-917840 p	UDP-N-acetylglucosamine 1-carboxyvinyltransferase
SEQ ID n° 6544	4408	2271	SA-917.1	918108-918965 p	similar to competence associated membrane nuclease
SEQ ID n° 6545	4409	2272	SA-918.1	919256-920296 p	Phenylalanyl-tRNA synthetase alpha chain
SEQ ID n° 6546	4410	2273	SA-919.1	920379-920900 p	similar to other proteins
SEQ ID n° 6547	4411	2274	SA-920.1	920954-923359 p	Phenylalanyl-tRNA synthetase beta chain
SEQ ID n° 6548	4412	2275	SA-921.1	923428-924096 m	similar to unknown proteins (C-terminal part)
SEQ ID n° 6549	4413	2276	SA-923.1	924207-927440 p	similar to ATP-dependent exonuclease, subunit B
SEQ ID n° 6550	4414	2277	SA-925.1	927430-931053 p	similar to ATP-dependent exonuclease, subunit A
SEQ ID n° 6551	4415	2278	SA-926.1	931066-931992 p	similar to cation transporter
SEQ ID n° 6552	4416	2279	SA-928.2	931967-933343 m	similar to putative tRNA modification GTPase TrmE
SEQ ID n° 6553	4417	2280	SA-93.1	2186362-2187234 p	similar to L-serine dehydratase alpha subunit
SEQ ID n° 6554	4418	2281	SA-930.2	866828-867787 m	Similar to ribonucleotide diphosphate reductase small subunit
SEQ ID n° 6555	4419	2282	SA-931.2	867990-870149 m	Similar to ribonucleotide reductase large subunit
SEQ ID n° 6556	4420	2283	SA-932.2	870227-870451 m	Similar to glutaredoxin
SEQ ID n° 6557	4421	2284	SA-934.1	870833-871096 p	Similar to histidine-containing phosphocarrier protein of the phosphotransferase system (PTS)
					phosphoenolpyruvate:sugar phosphotransferase system enzyme
SEQ ID n° 6558	4422	2285	SA-936.1	871101-872834 p	
SEQ ID n° 6559	4423	2286	SA-937.1	872984-874411 p	similar to glyceraldehyde-3-phosphate dehydrogenase
اء°	4424	2287	SA-939.1	874551-875804 p	similar to oligosaccharide deacetylase
SEQ ID n° 6561	4425	2288	SA-94.1	2187369-2188046 m	similar to other proteins



SEQ ID n° 6562	4426	2289	SA-941.1	875835-876917 m	similar to RNA helicase
SEQ ID n° 6563	4427	2290	SA-942.1	877062-877691 p	similar to uridine kinase
SEQ ID n° 6564	4428	2291	SA-943.1	877778-878275 p	similar to unknown proteins
SEQ ID n° 6565	4429	2292	SA-944.1	878275-879939 p	similar to DNA polymerase III gamma/tau subunit
SEQ ID n° 6566	4430	2293	SA-945.1	880028-880222 p	similar to unknown proteins
SEQ ID n° 6567	4431	2294	SA-946.1	880203-881138 m	similar to transcriptional repressor of the biotin operon
SEQ ID n° 6568	4432	2295	SA-947.1	881323-882519 p	S-adenosylmethionine synthetase
SEQ ID n° 6569	4433	2296	SA-948.1	883038-884945 p	similar to fibronectin-binding protein
SEQ ID n° 6570	4434	2297	SA-949.1	885012-885557 p	Unknown
SEQ ID n° 6571	4435	2298	SA-950.1	885717-885845 p	hypothetical protein
SEQ ID n° 6572	4436	2299	SA-951.1	885953-886519 p	similar to unknown proteins
SEQ ID n° 6573	4437	2300	SA-952.1	886516-887070 p	similar to unknown proteins
SEQ ID n° 6574	4438	2301	SA-953.2	887074-888360 p	similar to cation ABC transporter (ATP-binding protein)
SEQ ID n° 6575	4439	2302	SA-954.1	1170326-1171621 m	Unknown
SEQ ID n° 6576	4440	2303	SA-955.1	1171761-1172060 m	Similar to unknown protein
SEQ ID n° 6577	4441	2304	SA-956.1	1172071-1173318 m	Similar to DNA-methyltransferase
SEQ ID n° 6578	4442	2305	SA-957.1	1173315-1174946 m	Similar to plasmid relaxase and mobilisation protein A
SEQ ID n° 6579	4443	2306	SA-958.1	1174918-1175292 m	similar to unknown proteins
SEQ 1D n° 6580	4444	2307	SA-959.1	1175295-1175858 m	Unknown
					similar to aggregation promoting protein (adhesin involved in
SEQ ID n° 6581	4445	2308	SA-96.1	2188179-2188718 m	high frequency of conjugation)
SEQ ID n° 6582	4446	2309	SA-960.1	1175861-1176157 m	hypothetical CDS
SEQ ID n° 6583	4447	2310	SA-961.1	1176201-1176500 m	Unknown
SEQ ID n° 6584	4448	2311	SA-964.1	1176552-1179788 m	similar to plasmid unknown proteins
SEQ ID n° 6585	4449	2312	SA-966.1	1179790-1180155 m	Unknown
i i					Similar to transfer complex protein TrsK - Lactococcus lactis
SEQ 10 n° 6586	4450	2313	SA-967.1	1180197-1182242 m	plasmid pMRC01
SEQ ID n° 6587	4451	2314	SA-968.1	1182242-1182733 m	Unknown
SEQ ID n° 6588	4452	2315	SA-969.1	1182755-1183537 m	Similar to gram positive plasmid protein
SEQ ID n° 6589	4453	2316	SA-970.1	1183537-1183809 m	Unknown
SEQ ID n° 6590	4454	2317	SA-971.1	1183829-1184434 m	Unknown
SEQ ID n° 6591	4455	2318	SA-973.1	1184455-1187145 m	similar to plasmid unknown proteins
SEQ ID n° 6592	4456	2319	SA-974.1	1187178-1187708 p	Unknown
SEQ ID n° 6593	4457	2320	SA-975.2	1187902-1190292 m	Similar to plasmid transfer complex protein TrsE
SEQ ID n° 6594	4458	2321	SA-976.2	1899710-1900897 m	similar to two-component sensor histidine kinase
SEQ ID n° 6595	4459	2322	SA-977.1	1898029-1899564 m	Similar to D-alanine-D-alanyl carrier protein ligase

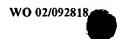


SEQ ID n° 6596	4460	2323	SA-978.1	1896767-1898032 m	1896767-1898032 m   Similar to LTA D-alanylation protein DItB
SEQ 1D n° 6597	4461	2324	SA-98.1	2188934-2189728 m	putative ABC transporter (permease)
SEQ ID n° 6598	4462	2325	SA-980.1	1896513-1896752 m	Similar to D-alanyl carrier protein
SEQ ID n° 6599	4463	2326	SA-981.1	1895258-1896520 m	similar to LTA D-alanine transfer protein DItD
SEQ ID n° 6600	4464	2327	SA-982.1	1894691-1895083 m	Unknown
SEQ ID n° 6601	4465	2328	SA-983.1	1893399-1894691 m	Unknown .
SEQ ID n° 6602	4466	2329	SA-984.1	1893016-1893408 m	Unknown
SEQ ID n° 6603	4467	2330	SA-985.1	1892725-1893006 m	Unknown
SEQ ID n° 6604	4468	2331	SA-986.4	1891761-1892543 m	similar to unknown proteins
SEQ ID n° 6605	4469	2332	SA-987.4	1891208-1891774 m	similar to unknown proteins
SEQ ID n° 6606	4470	2333	SA-988.4	1890755-1891204 m	histidine triad family protein
SEQ ID n° 6607	1471	2334	SA-989.1	1889856-1890728 m	Similar to 16 rRNA (adenine-N6,N6-)-dimethyltransferase
SEQ ID n° 6608	4472	2335	SA-99.1	2189721-2190563 m	1.2 Transport/binding proteins and lipoproteins
SEQ ID n° 6609	4473	2336	SA-990.1	1889001-1889852 m	Similar to 23S ribosomal RNA methyltransferase
SEQ ID n° 6610	4444	2337	SA-991.1	1887973-1888845 m	similar to unknown proteins
SEQ ID n° 6611	4475	2338	SA-993.1	1887304-1887966 m	similar to ribulose-5-phosphate 3-epimerase
SEQ ID n° 6612	9446	2339	SA-994.1	1886679-1887311 m	Similar to unknown proteins
SEQ ID n° 6613	4477	2340	SA-995.1	1885403-1886677 m	Similar to unknown proteins
SEQ ID n° 6614	4478	2341	SA-996.1	1884472-1885413 m	similar to unknown proteins
SEQ ID n° 6615	6444	2342	SA-997.2	1883563-1884375 m	similar to transcription repressor of purine operon PurR
SEQ ID n° 6616	4480	2343	SA-998.2	200712-201377 p	qimilar to unknown proteins
SEQ ID n° 6617	4481	2344	SA-999.1	201398-202168 m	similar to pyrroline-5-carboxylate reductase



<u>TABLEAU 4</u>. Localisation des 139 contigs de séquence SEQ ID No. 1 à SEQ ID No. 139 sur la séquence génomique complète (SEQ ID No. 2345).

		Position sur complet (= SEC		Sens
SEQ ID	Contig	position Début	position Fin	(m=minus / p=plus)
SEQ ID n°1	Contig1	1356648	1355991	m
SEQ ID n°2	Contig2	341651	341120	m
SEQ ID n°3	Contig3	476798	476418	m
SEQ ID n°4	Contig4	1475712	1475086	m
SEQ ID n°5	Contig5	784818	784243	m
SEQ ID n°6	Contig6	1756826	1757251	р
SEQ ID n°7	Contig7	1950108	1949648	m
SEQ ID n°8	Contig8	138225	138876	р
SEQ ID n°9	Contig9	2097891	2098283	р
SEQ ID n°10	Contig10	1238491	1237984	m
SEQ ID n°11	Contig11	1882130	1881745	m
SEQ ID n°12	Contig12	1089348	1088935	m
SEQ ID n°13	Contig13	555788	555189	m
SEQ ID n°14	Contig14	2017928	2017437	m ·
SEQ ID n°15	Contig15	1154094	1154701	р
SEQ ID n°16	Contig16	752647	753091	р
SEQ ID n°17	Contig17	1355561	1355078	m
SEQ ID n°18	Contig18	1255951	1256101	р
SEQ ID n°19	Contig19	792712	793148	р
SEQ ID n°20	Contig20	481787	482228	P
SEQ ID n°22	Contig22	1590263	1590842	р
SEQ ID n°23	Contig23	508269	508918	P
SEQ ID n°24	Contig24	1142198	1142488	р
SEQ ID n°25	Contig25	1982019	1981737	m
SEQ ID n°26	Contig26	119342	119919 .	P
SEQ ID n°28	Contig28	1124069	1123256	m
SEQ ID n°29	Contig29	266586	266900	р
SEQ ID n°30	Contig30	111013	111623	Р
SEQ ID n°31	Contig31	1804173	1804706	р
SEQ ID n°32	Contig32	2170341	2169828	m
SEQ ID n°33	Contig33	1959867	1959394	m ·
SEQ ID n°34	Contig34	1295529	1294939	·m
SEQ ID n°35	Contig35	178592	178071	m
SEQ ID n°36	Contig36	1857103	1856614	m
SEQ ID n°37	Contig37	1063484	1063911	р
SEQ ID n°38	Contig38	198025	197570	m
SEQ ID n°39	Contig39	1486076	1486553	р
SEQ ID n°40	Contig40	2033914	2034352	р
SEQ ID n°41	Contig41	737932	738486	Р
SEQ ID n°42	Contig42	729008	728453	m
SEQ ID n°43	Contig43	1671733	1672151	р



SEQ ID n°44	Contig44	1103091	1103644	р
SEQ ID n°45	Contig45	700139	699583	m
SEQ ID n°46	Contig46	207521	206897	m
SEQ ID n°47	Contig47	1064808	1065099	р
SEQ ID n°48	Contig48	1091636	1092281	P
SEQ ID n°49	Contig49	1701764	1700906	m
SEQ ID n°50	Contig50	609072	609590	р
SEQ ID n°51	Contig51	1459271	1458780	m
SEQ ID n°52	Contig52	60603	60154	m
SEQ ID n°53	Contig53	289646	289284	m
SEQ ID n°54	Contig54	1536438	1536058	m
SEQ ID n°55	Contig55	509420	510430	р
SEQ ID n°56	Contig56	1559964	1558709	· m
SEQ ID n°58	Contig58	2166712	2165923	m
SEQ ID n°59	Contig59	1919605	1920984	р
SEQ ID n°60	Contig60	962333	960438	m
SEQ ID n°61	Contig61	1363649	1365724	р
SEQ ID n°62	Contig62	1140306	1137284	m
SEQ ID n°63	Contig63	1702242	1706039	Р
SEQ ID n°64	Contig64	1490271	1493283	р
SEQ ID n°65	Contig65	783206	785628	р
SEQ ID n°66	Contig66	852318	849615	m
SEQ ID n°67	Contig67	1882303	1880181	m
SEQ ID n°68	Contig68	1614050	1618058	р
SEQ ID n°69	Contig69	1484885	1490042	р
SEQ ID n°70	Contig70	510495	516449	р
SEQ ID n°71	Contig71	125082	121213	m
SEQ ID n°72	Contig72	1557644	1551892	m
SEQ ID n°73	Contig73	145707	143269	m
SEQ ID n°74	Contig74	859105	852465	m
SEQ ID n°75	Contig75	1219383	1215342	m
SEQ ID n°76	Contig76	1091627	1086724	m
SEQ ID n°77	Contig77	1245975	1251984	р
SEQ ID n°78	Contig78	115260	121688	Р
SEQ ID n°79	Contig79	1100300	1092624	m
SEQ ID n°80	Contig80	1107948	1100525	m
SEQ ID n°81	Contig81	1245466	1237461	m
SEQ ID n°82	Contig82	2111296	2104033	m
SEQ ID n°83	Contig83	33479	27132	m
SEQ ID n°84	Contig84	1339614	1350526	р
SEQ ID n°85	Contig85	2070423	2058143	m
SEQ ID n°86	Contig86	1462530	1470059	р
SEQ ID n°87	Contig87	526582	517432	m
SEQ ID n°88	Contig88	1484487	1470171	m
SEQ ID n°89	Contig89	1124087	1136746	p
SEQ ID n°90	Contig90	1879890	1866931	m
SEQ ID n°91	Contig91	1721684	1706045	m
SEQ ID n°92	Contig92	1358184	1357897	æ
SEQ ID n°93	Contig93	1577596	1560798	m
SEQ ID n°94	Contig94	115130	103188	m



SEQ ID n°95	Contig95	1921051	1933881	р
SEQ ID n°96	Contig96	1944905	1933782	m
SEQ ID n°97	Contig97	1919624	1906953	, m
SEQ ID n°98	Contig98	2090559	2103658	р
SEQ ID n°99	Contig99	1237482	1219423	m
SEQ ID n°100	Contig100	1123110	1108191	m
SEQ ID n°101	Contig101	1551836	1529458	m
SEQ ID n°102	Contig102	1818811	1800978	m
SEQ ID n°103	Contig103	764781	783195	р
SEQ ID n°104	Contig104	1086606	1065938	m
SEQ ID n°105	Contig105	125425	143102	р
SEQ ID n°106	Contig106	962438	984387	р
SEQ ID n°107	Contig107	1169838	1190193	р
SEQ ID n°108	Contig 108	2090426	2070667	m
SEQ ID n°109	Contig109	1140315	1169462	р
SEQ ID n°111	Contig111	238297	258413	р
SEQ ID n°112	Contig112	216686	237881	P
SEQ ID n°113	Contig113	2209521	16967	m
SEQ ID n°114	Contig114	1883537	1906918	р
SEQ ID n°115	Contig115	145772	172009	р
SEQ ID n°116	Contig116	508181	477405	m
SEQ ID n°117	Contig117	859233	888273	р
SEQ ID n°118	Contig118	1529046	1494213	m
SEQ ID n°119	Contig119	473132	438871	æ
SEQ ID n°120	Contig120	1981657	1945366	m
SEQ ID n°121	Contig121	1613824	1577594	m
SEQ ID n°122	Contig122	1765846	1800817	Р
SEQ ID n°123	Contig123	2111499	2153851	р
SEQ ID n°124	Contig124	1721668	1765765	p
SEQ ID n°125	Contig125	984406	1025178	Р
SEQ ID n°126	Contig126	1293488	1339586	ρ
SEQ ID n°127	Contig127	216691	176332	m
SEQ ID n°128	Contig128	1818941	1866861	Р
SEQ ID n°129	Contig129	849565	785796	m
SEQ ID n°130	Contig130	888292	960270	р
SEQ ID n°131	Contig131	2208563	2155215	m
SEQ ID n°132	Contig132	33590	88257	р
SEQ ID n°133	Contig133	1982609	2057812	Р
SEQ ID n°134	Contig134	1700642	1618142	m
SEQ ID n°135	Contig135	1293063	1190375	m
SEQ ID n°136	Contig136	1366980	1462324	р
SEQ ID n°137	Contig137	390853	434186	р
SEQ ID n°138	Contig138	357393	259739	m
SEQ ID n°139	Contig139	527049	716899	р

<u>TABLEAU 5</u>. Propriété d'adhérence à des cellules épithéliales humaines en culture de la souche NEM316 de *S. agalactiae* et de souches mutantes dérivées.

Souche	Gène inactivé	% d'adhésion <sup>a</sup>		
		Cellules A549	Cellules Hela	
NEM316	aucun	9	16	
NEM1979	IPF N° 1268 (srtA)	1,5	2	
NEM2056	IPF N° 678	. 2	n.t.	
NEM2057	IPF N° 1503	4,5	n.t.	

5 a, le pourcentage d'adhésion correspond au nombre de bactéries (Unité Formant des Colonies, UFC) restant adhérentes aux cellules après lavage avec du tampon PBS par rapport au nombre d'UFC ajoutées à la monocouche de cellules épithéliales.



TABLEAU 6. Gènes de la souche de S. agalactiae NEM316 codant pour des protéines de surface avec un motif d'ancrage LPXTGª

Fonction prédite <sup>c</sup>	inconnue	Protéine liant l'acide sialique	Protéine liant l'acide sialique	Adhésine	inconnue Adhésine	inconnue	inconnue	inconnue Protéine liant la choline	Serine proteinase	Cyclo-nucleotide phosphodiesterase
% d'acides aminé identiques (similaires) / longueur de la région similaire	74 (77) / 798 71 (76) / 877 69 (75) / 1103	50 (60) / 1314 43 (53) / 1248	30 (43) / 1385 31 (45) / 1285	25 (38) / 358	31 (46) / 302 23 (38) / 795	38(52) / 406		50(62) / 183 30(60) / 220	49 (65) / 1596	57(70) / 694 47(66) / 630
Protéines homologues	Alp2 (S. agalactiae) Alp3 (S. agalactiae) R28 (S. pyogenes)	Hsa (S. gordonii) SrpA (S. cristatus)	Ssp-5 (S. gordonii), PAa (S. intermedius)	EaeH (E. coli O157:H7)	M-like protein (S. equi) PspC (S. pneumoniae)	SpaA (S. sobrinus)	No homology in public databases	Cell surface protein (S. mutans) CbpD (S. pneumoniae)	PrtS (S. thermophilus)	CpdB (S. dysgalactiae) YfkN (Bacillus subtilis)
Site de coupure	LPXT/G	=	=	=	£	=	=	z	=	:
taille pbases	1126	1310	1634	512	643	932	308	543	1570	800
IPF N°	523	571	220	2192	1716	1247	2337	1861	1503	678
Seq ID (ADN)	6194	6236	5497	5491	5103	4705	2610	5234	4926	6331

										417									
( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( )	Amindase	Amylopullulanase alkaline	inconnue		inconnue	inconnue	inconnue	inconnue	Surface exclusion protein	inconnue	Surface exclusion pretein	inconnue	inconnue	inconnue	Protéine liant la fibronectine	inconnue	inconnue	inconnue	inconnue
36 (54) / 478	35 (54) / 492	65 (79) / 1095	23(47) / 373	,	26(50) / 273	37(52) / 405	36(52) / 399	33 (49) / 225	24 (37) / 715	31 (47) / 263	22 (40) / 784	33 (47) / 211			32(46) / 176	27(42) / 512	25(38) / 577		
AmiC (S. pyogenes)	YbgE (L. lactis)	PulA (S. pyogenes)	CG15040 gene product Drosophila	melanogaster	Antigen p200 (Babesia bigemina)	SpaA (S. sobrinus)	Pas (S. intermedius)	Plasmid-encoded protein (E. faecalis)	Sec10 (E. faecalis)	Plasmid-encoded protein (E. faecalis)	Sec10 (E. faecalis)	Plasmid-encoded protein (E. faecalis)	No homology in public databases	No homology in public databases	PFBP (S. pyogenes)	Hypothetical protein 2 (Lactobacillus leichmannii)	Fimbrial structural subunit (Actinomyces naeslundii)	No homology in public databases	No homology in public databases
	•	=		=		=		=	=	z.	=	Ξ	=	E	IPXT/G	Ξ		=	
089	}	1252		410		033	3	240	753	236	743	253	192	521	901	733	<b>t</b>	307	674
585	)	280		2495		15	:	11	18	1248	1250	2414	584*	765	2300	908		807	2334
6247		5842		5741		4921		2090	5180	4706	4708	2295	6246	6411	5578	6446		6447	2607



		1	T	$\top$	
inconnue	inconnue	Protéase à sérine	C5a peptidase	5'-nucleotidase secretée	inconnue
72 (81) / 1050	24 (41) / 566	38(55) / 1194	99(99) / 1150	60(74) / 688	
SPy0843 (S. pyogenes)	BspA (Bacteroides forsythus)	ScpB (S. agalactiae)	ScpB (S. agalactiae)	SPy0872 (S. pyogenes)	No homology in public databases
S/LXd I		=	LPXT/N	z	FPKT/G
1055		1233	1150	069	068
538	) )	547	2082	2390	1551
6209		,6215	5406	5658	4965

a, Les protéines ancrées au peptidoglycane ont été identifiées par la recherche d'un motif LPXTG ou un motif voisin C-terminal suivi d'un domaine hydrophobe et d'acides aminés basiques. Les similarités par BLASTP avec des protéines à domaine LPXTG connues ont aussi été utilisées. b, Seulement les similarité avec une probabilité BLASTP <10-10 ont été considérées comme significatives. C, La fonction a été prédite par analogie avec celles des protéines homologues contenues dans la bases de séquences protéique nrprot du NCBI.

<u>TABLEAU 7</u>. Distribution des gènes codant pour des protéines de surface à motif LPXTG parmi des isolats cliniques indépendants de 5 sérotypes de S. agalactiae.

Seq ID	N° d'IPF		Proportion des souches portant le gène							
(ADN)		Type Ia 23 isolats	Type Ib 7 isolats	Type II 12 isolats	Type III 39 isolats	Type V 16 isolats	non groupé 2 isolats	Fréquence totale 99 isolats		
4926	1503	100	100	100	100	100	100	100		
6331	678	100	100	100	100	100	100	100		
5491	2192	100	100	100	100	100	100	100		
5234	1861	100	100	100	100	100	100	100		
6246	584	100	100	100	100	100	100	100		
5842	280	100	100	100	100	100	100	100		
6247	585	86,9	85,7	91,7	92,3	81,3	100	88,9		
4965	1551	34,8	85,7	100	94,8	93,8	100	80,8		
6447	807	26,1	85,7	83,3	92,3	93,8	100	79,8		
5610	2337	86,9	100	100	56,4	100	50	78,8		
6236	571	82,6	100	83,3	46,1	100	50	71,7		
5103	1716	91,0	100	83,3	38,5	81,3	50	67,7		
5607	2334	56,5	100	83,3	46,1	100	50	65,6		
6411	765	30,4	71,4	41,7	61,5	87,5	100	57,6		
4921	15	65,2	0	41,7	59	12,5	50	46,5		
5090	17	4,3	14,3	16,7	20,5	6,3	0	13,1		
5180	18	4,3	14,3	0	20,5	6,3	0	11,1		
4706	1248	0	0	0	12,8	6,3	0	6		
5497	220	0	14,3	8,3	5,1	0	0	4,5		
4708	1250	0	14,3	0	5,1	0	0	3		
5677	2414	0	0	0	7,7	0	0	3		



## TABLEAU 8. Lipoprotéines

6327 9 unknown 6330 339 similar to unknown proteins 635 344 Similar to ABC transporter (binding protein) 6137 460 Similar to ABC transporter (binding protein) 6138 504 similar to unknown proteins 6294 638 Similar to unknown proteins 6335 682 Similar to oligopeptide and pheromone binding protein 6386 739 similar to other lipoprotein 6386 739 similar to (oligopeptide) ABC transporter (binding protein) 4495 1018 Similar to (oligopeptide) ABC transporter (binding protein) 4596 1119 similar to ribose ABC transporter (binding protein) 4636 1162 similar to (amino acid?) ABC transporter (binding protein) 4730 1280 similar to ABC transporter (binding protein) 4836 1399 similar to phosphate ABC transporter (binding protein) 4836 1399 similar to phosphate ABC transporter (binding protein) 4906 1481 Similar to D,D-carboxypeptidase 4920 1499 similar to peptidyl-prolyl cis-trans isomerase 4925 1502 similar to metal ABC transporter (binding protein) 5021 1617 Similar to unknown lipoprotein 5158 1775 similar to oligopeptide ABC transporter (binding protein) 5247 1879 similar to oligopeptide ABC transporter (binding protein) 5267 1879 similar to glycine betaine/carnitine/choline ABC transporter (osmoprotec	
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5021 1617 Similar to unknown lipoprotein  5158 1775 similar to ferrichrome ABC transporter (binding protein)  5247 1879 similar to oligopeptide ABC transporter (binding protein)  similar to glycine betaine/carnitine/choline ABC transporter (osmoprotes)	
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5247 1879 similar to oligopeptide ABC transporter (binding protein)  similar to glycine betaine/carnitine/choline ABC transporter (osmoprotes)	
similar to glycine betaine/carnitine/choline ABC transporter (osmoprotec	
similar to glycine betaine/carnitine/choline ABC transporter (osmoprotec	
5306   1955   binding protein)	tant-
5417 2099 similar to putative ABC transporter (binding protein)	
5423 2103 Unknwon, similar to unknown protein and to B. subtilis SpoIIIJ protein	
5450 2133 laminin-binding surface protein	
5486 2185 putative ABC transporter (binding protein)	

5559	2278	putative ABC transporter (binding protein)
5591	2314	similar to protease maturation protein
5677	2414	Similar to plasmid related proteins, Putative peptidoglycan bound protein (LPXTG motif)
5718	2464	Similar to unknown proteins
5732	2482	similar to ferrichrome ABC transporter (binding protein)
5799	2597	similar to amino acid ABC transporter (binding protein)
5800	2598	similar to phosphate ABC transporter (binding protein)
5837	2789	Unknown
5861	2843	Similar to amino acid ABC transporter (binding protein)
5883	2875	Unknown
59:23	2922	Similar to amino acid ABC transporter (binding protein)

Les gènes codant pour des lipoprotéines ont été identifiés sur la base de la prédiction du motif de coupure / modification de type lipoprotéine [S. Hayashi, H. C. Wu. J Bioenerg Biomembr. 22, 451 (1990)] et d'un peptide signal (identifié en utilisant SignalP vs2.0 [H Nielsen, *Prot Engin* 12, 13-9. (1999)]) et par l'analyse des résultats de comparaison sur les banques de séquence protéiques en utilisant BLAST [S. F. Altschul et al., *Nucleic Acids Res* 25, 3389-402. (1997)].

Tableau 9. Autres protéines de surface

Seq ID (ADN)	Nº d'IPF	Annotation
6037	347	group B streptococcal surface immunogenic protein
4972	1562	Putative cell wall protein, weakly similar to peptidase or esterase
6569	948	similar to fibronectin-binding protein
5234	1861	similar to cell wall proteins
5530	2238	possible surface protein
5223	1847	CAMP factor



Ces protéines ont été identifiées sur la base de similarité avec d'autres protéines de surface bactérienne et la prédiction d'un peptide signal et ne faisant pas partie des classes des protéines ancrées au peptidoglycane et lipoprotéines.

5 <u>TABLEAU 10</u>. Protéines impliquées dans la biosynthèse de composés polysaccharidiques de la paroi de S. agalactiae.

Seq ID (ADN)	N° d'IPF	Annotation	Commentaires
4861	1430	similar to Streptococcus mutans RgpG protein required for	
4001	1450	biosynthesis of rhamnose-glucose polysaccharide	
6214	544	similar to rhamnosyltransferase	
6061	381	Unnown, Similar to UDP-N-acetylmuramoylalanineD-	
0001	361	glutamate ligase	
6517	890	similar to Cell Wall Muropeptide Branching Enzyme	
6518	891	similar to cell wall muropeptide branching enzyme	
6519	892	similar to cell wall muropeptide branching enzyme	
4743	1295	similar to glycosyltransferases	
6343	692	similar to hexosyltransferase	
6342	691	similar to glucosyl transferase	
5326	1977	Similar to UDP-D-glucose:galactosyl glucosyltransferase	
4952	1532	similar to N-acetylneuraminic acid synthetase	
5619	2346	capsular polysaccharide biosynthesis protein	1
5618	2345	similar to glycosyl transferase	
5617	2344	similar to glycosyl transferase	Biosynthèse de
5616	2343	capsular polysaccharide repeating-unit polymerase	la capsule
5615	2342	beta-1,4-galactosyltransferase	
5614	2341	beta-1,4-galactosyltransferase enhancer	
5613	2340	similar to glucose-1-phosphate transferase	
5611	2339	capsular polysaccharide chain length regulator/exporter	
5696	2437	putative chain length regulator CpsC	
5971	301	similar to dTDP-glucose-4,6-dehydratase	
5233	1860	similar to to Cell Wall Muropeptide Branching Enzyme	

5602	2220		
3602	2329	similar to capsular polyglutamate biosynthesis	
5156	1773	Similar to UDP-N-acetylmuramoylalanyl-D-glutamyl-2, 6-	
3150	1775	diaminopimelate ligase	
5574	2297	Similar to putative hexosyltransferase	
5573	2296	Similar to rhamnosyl transferase I	
5654	2386	Similar to capsular polysaccharide synthesis protein	
5656	2388	Similar to putative rhamnosyltransferase	
5526	2233	Similar to putative rhamnosyltransferase	
5527	2235	Similar to nucleotide-sugar dehydratase	
5529	2237	Similar to Pneumococcal LicD2 protein involved in	
3327	2231	phosphorylcholine metabolism	
5534	2241	similar to rhamnosyltransferase	
5625	2354	similar to putative rhamnosyltransferase	
5626	2355	dTDP-L-rhamnose synthase	
6223	555	Similar to putative glucosyl transferase	
6229	562	Similar to hypothetical glycosyl transferase	
6230	563	Similar to putative glycosyltransferase	
6231	565	Similar to putative glycosyl transferase	
6232	566	Similar to putative glycosyl transferase	
6233	567	Similar to putative glycosyl transferase	1 "" 7 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
5764	2518	similar to putative sugar transferase	
6095	416	similar to UDP-N-acetylglucosamine pyrophosphorylase	
5089	1699	Similar to UDP-N-acetylmuramate-alanine ligase	
5466	2158	similar to glycosyl transferase	
5465	2157	similar to glycosyl transferase	

Ces gènes ont été identifiés par l'analyse des résultats de similarité avec les séquences protéiques connues en utilisant le logiciel BLASTP. Les produits de ces gènes pourraient intervenir dans la biosynthèse de polysaccharides qui pourraient être des constituants de préparations vaccinales.

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## REVENDICATIONS

- 1. Séquence nucléotidique isolée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi les séquences SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345.
- 2. Séquence nucléotidique isolée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi :
- a) une séquence nucléotidique comportant au moins 75 % d'identité avec une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345;
- b) une séquence nucléotidique hybridant dans des conditions de forte stringence avec une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, et comprenant au moins 20 nucléotides;
  - c) une séquence nucléotidique complémentaire d'une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, ou complémentaire d'une séquence nucléotidique telle que définie en a), ou b), ou une séquence nucléotidique de l'ARN correspondant à l'une des séquences a) ou b);
  - d) une séquence nucléotidique d'un fragment représentatif d'une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, ou d'un fragment représentatif d'une séquence nucléotidique telle que définie en a), b) ou c) et comprenant au moins 20 nucléotides;
  - e) une séquence nucléotidique comprenant une séquence telle que définie en a), b), c) ou d); et
  - f) une séquence nucléotidique telle que définie en a), b), c), d) ou e) modifiée et comportant au plus 10 % de nucléotides modifiés par rapport à la séquence de référence.
  - 3. Séquence nucléotidique selon la revendication 2, caractérisée en ce qu'il s'agit d'une séquence issue d'une séquence choisie parmi SEQ ID No. 1 à SEQ ID No. 139 et SEQ ID No. 2345, et en ce qu'elle code pour un polypeptide, choisi de préférence parmi les séquences SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481.
  - 4. Séquence nucléotidique caractérisée en ce qu'elle comprend une séquence nucléotidique choisie parmi :
  - a) une séquence nucléotidique selon la revendication 3 ou choisie parmi les séquences SEQ ID No. 4482 à SEQ ID No. 6617;

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- b) une séquence nucléotidique comportant au moins 75 % d'identité avec une séquence nucléotidique selon la revendication 3 ;
- c) une séquence nucléotidique s'hybridant dans des conditions de forte stringence avec une séquence nucléotidique selon la revendication 3 et comprenant au moins 20 nucléotides;
- d) une séquence nucléotidique complémentaire ou d'ARN correspondant à une séquence telle que définie en a), b) ou c);
- e) une séquence nucléotidique d'un fragment représentatif d'une séquence telle que définie en a), b), c) ou d) et comprenant au moins 20 nucléotides ; et
- f) une séquence telle que définie en a), b), c), d) ou e) modifiée et comportant au plus 10 % de nucléotides modifiés par rapport à la séquence de référence.
  - 5. Polypeptide codé par une séquence nucléotidique selon l'une des revendications 2 à 4.
  - Polypeptide selon la revendication 5, caractérisé en ce qu'il est choisi parmi les polypeptides choisis parmi SEQ ID No. 140 à SEQ ID No. 2344, et SEQ ID No. 2346 à SEQ ID No. 4481.
    - 7. Polypeptide caractérisé en ce qu'il comprend un polypeptide choisi parmi :
    - a) un polypeptide selon l'une des revendications 5 et 6;
    - b) un polypeptide présentant au moins 80 % d'identité avec un polypeptide selon l'une des revendications 5 et 6;
    - c) un fragment d'au moins 5 acides aminés d'un polypeptide selon l'une des revendications 5 et 6, ou tel que défini en b);
    - d) un fragment biologiquement actif d'un polypeptide selon l'une des revendications 5 et 6, ou tel que défini en b) ou c); et
- e) un polypeptide selon l'une des revendications 5 et 6 ou tel que défini en b), c) ou d) modifié et comportant au plus 10 % d'acides aminés modifiés par rapport à la séquence de référence.
  - 8. Séquence nucléotidique codant pour un polypeptide selon la revendication
- 9. Séquence nucléotidique isolée codant pour un polypeptide spécifique de Streptococcus agalactiae choisi parmi les polypeptides de séquence SEQ ID No. 140 à SEQ ID No. 2344 et SEQ ID No. 2346 à SEQ ID No. 4481.

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- 10. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans la biosynthèse des acides aminés ou l'un de ses fragments.
- 11. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans la biosynthèse des cofacteurs, groupes prosthétiques et transporteurs ou l'un de ses fragments.
- 12. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide d'enveloppe cellulaire ou situé à la surface de *Streptococcus agalactiae* ou l'un de ses fragments.
- 13. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans la machinerie cellulaire ou l'un de ses fragments.
- 14. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme intermédiaire central ou l'un de ses fragments.
- 15. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme énergénique ou l'un de ses fragments.
- 16. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme des acides gras et des phospholipides ou l'un de ses fragments.
- 17. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme des nucléotides, des purines, des pyrimidines ou nucléosides ou l'un de ses fragments.
- 18. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans les fonctions de régulation ou l'un de ses fragments.
- 19. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de réplication ou l'un de ses fragments.

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- 20. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de transcription ou l'un de ses fragments.
- 21. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de traduction ou l'un de ses fragments.
- 22. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de transport et de liaison des protéines ou l'un de ses fragments.
- 23. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans l'adaptation aux conditions atypiques ou l'un de ses fragments.
- 24. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans la sensibilité aux médicaments et analogues ou l'un de ses fragments.
- 25. Séquence nucléotidique selon l'une des revendications 2 à 4, 8 et 9, caractérisée en ce qu'elle code pour un polypeptide de *Streptococcus agalactiae* impliqué dans les fonctions relatives aux transposons ou l'un de ses fragments.
- 26. Séquence nucléotidique isolée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi :
  - a) une séquence choisie parmi les séquences SEQ ID N° 6194,6236,5497,5791,5103,4705,5610,5234,4926,6331,6247,5842,5741,4921,5090,518 0,4706,4708,5677,6246,6411,5578,6446,6447,5607,6209,6215,5406,5658,4965, de préférence parmi les séquences SEQ ID N°4926,6331,5491,5234,6246,5842;
  - b) une séquence nucléotidique comportant au moins 75 % d'identité avec une séquence nucléotidique du a);
  - c) une séquence nucléotidique s'hybridant dans des conditions de forte stringence avec une séquence nucléotidique du a) ou b) et comprenant au moins 20 nucléotides;
  - d) une séquence nucléotidique complémentaire ou d'ARN correspondant à une séquence telle que définie en a), b) ou c);
  - e) une séquence nucléotidique d'un fragment représentatif d'une séquence telle que définie en a), b), c) ou d) et comprenant au moins 20 nucléotides; et

f) une séquence telle que définie en a), b), c), d) ou e) modifiée et comportant au plus 10 % de nucléotides modifiés par rapport à la séquence de référence ;

et en ce qu'elle code pour une protéine de surface avec un motif d'ancrage LPXTG.

- 5 27. Séquence nucléotidique isolée de *Streptococcus agalactiae*, caractérisée en ce qu'elle est choisie parmi les séquences SEQ ID N°6035,6137,6335,6377,6386,4495,4596,4636,4730,4816,4836,4906,4920,4925,5158,5 247, 5306,5417,5450,5486,5559,5591,5677,5732,5799,5800,5861,5923 et en ce qu'elle code pour une lipoprotéine.
- 10 28. Séquence nucléotidique isolée de Streptococcus agalactiae, caractérisée en ce qu'elle choisie parmi les séquences SEQ  $\mathbf{m}$ N°4861,6214,6061,6517,6518,6519,4743,6343,6342,5326,4952,5619,5618,5617,5616,5 615,5614,5613,5611,5696,5971,5233,5602,5156,5574,5573,5654,5656,5526,5527,5529 ,5534,5625,5626,6223,6229,6230,6231,6232,6233,5764,6095,5089,5466,5465 et en ce qu'elle code pour une protéine impliquée dans la biosynthèse de composés 15 polysaccharidiques de paroi.
  - 29. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans la biosynthèse des acides aminés ou l'un de ses fragments.
- 30. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans la biosynthèse des cofacteurs, groupes prosthétiques et transporteurs ou l'un de ses fragments.
  - 31. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide d'enveloppe cellulaire ou situé à la surface de *Streptococcus* agalactiae ou l'un de ses fragments.
  - 32. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans la machinerie cellulaire ou l'un de ses fragments.
  - 33. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme intermédiaire central ou l'un de ses fragments.
  - 34. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme énergétique ou l'un de ses fragments.

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- 35. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme des acides gras et des phospholipides ou l'un de ses fragments.
- 36. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le métabolisme des nucléotides, des purines, des pyrimidines ou nucléosides ou l'un de ses fragments.
- 37. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans les fonctions de régulation ou l'un de ses fragments.
- 38. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de réplication ou l'un de ses fragments.
- 39. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de transcription ou l'un de ses fragments.
- 40. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de traduction ou l'un de ses fragments.
- 41. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans le processus de transport et de liaison des protéines ou l'un de ses fragments.
- 42. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans l'adaptation aux conditions atypiques ou l'un de ses fragments.
- 43. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans la sensibilité aux médicaments et analogues ou l'un de ses fragments.
  - 44. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il s'agit d'un polypeptide de *Streptococcus agalactiae* impliqué dans les fonctions relatives aux transposons ou l'un de ses fragments.
  - 45. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il est codé par une séquence selon la revendication 26 et est une protéine de surface avec un motif d'ancrage LPXTG.

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- 46. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il est codé par une séquence selon la revendication 27 et est une lipoprotéine.
- 47. Polypeptide selon l'une des revendications 5 à 7, caractérisé en ce qu'il est codé par une séquence selon la revendication 28 et est une protéine impliquée dans la biosynthèse de composés polysaccharidiques de paroi.
- 48. Séquence nucléotidique utilisable comme amorce ou comme sonde, caractérisée en ce que ladite séquence est choisie parmi les séquences nucléotidiques selon l'une des revendications 2 à 4, 8 à 28.
- 49. Séquence nucléotidique selon la revendication 48, caractérisée en ce qu'elle est marquée par un composé radioactif ou par un composé non radioactif.
- 50. Séquence nucléotidique selon l'une des revendications 48 et 49, caractérisée en ce qu'elle est immobilisée sur un support, de manière covalente ou non-covalente.
- 51. Séquence nucléotidique selon la revendication 50, caractérisée en ce qu'elle est immobilisée sur un support tel qu'un filtre à haute densité ou une puce à ADN.
- 52. Séquence nucléotidique selon l'une des revendications 49 à 51 pour la détection et/ou l'amplification de séquences nucléiques.
  - 53. Puce à ADN ou filtre, caractérisée en ce qu'elle contient au moins une séquence nucléotidique selon la revendication 51.
  - 54. Puce à ADN ou filtre selon la revendication 53, caractérisée en ce qu'elle contient en outre au moins une séquence nucléotidique d'un micro-organisme autre que Streptococcus agalactiae, immobilisée sur le support de ladite puce.
  - 55. Puce à ADN ou filtre selon la revendication 54, caractérisée en ce que le micro-organisme autre est choisi parmi un micro-organisme associé à *Streptococcus* agalactiae, une bactérie du genre *Streptococcus*, et un variant de *Streptococcus* agalactiae.
  - 56. Kit ou nécessaire pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé, caractérisé en ce qu'il comprend une puce à ADN ou un filtre selon la revendication 53.
  - 57. Kit ou nécessaire pour la détection et/ou l'identification d'un microorganisme, caractérisé en ce qu'il comprend une puce à ADN ou un filtre selon l'une des revendications 54 et 55.
    - 58. Kit ou nécessaire pour la détection et/ou la quantification de l'expression d'au moins un gène de *Streptococcus agalactiae*, caractérisé en ce qu'il comprend une puce à ADN ou un filtre selon l'une des revendications 53 à 55.

- 59. Vecteur de clonage, et/ou d'expression, caractérisé en ce qu'il contient une séquence nucléotidique selon l'une des revendications 1 à 4, 8 à 28.
- 60. Cellule hôte, caractérisée en ce qu'elle est transformée par un vecteur selon la revendication 59.
- 61. Cellule hôte selon la revendication 60, caractérisée en ce qu'il s'agit d'une bactérie appartenant au genre *Streptococcus*.
  - 62. Cellule hôte selon la revendication 61, caractérisée en ce qu'il s'agit d'une bactérie appartenant à l'espèce Streptococcus agalactiae.
- 63. Végétal ou animal, excepté l'Homme, comprenant une cellule transformée selon l'une des revendications 60 à 62.
  - 64. Procédé de préparation d'un polypeptide, caractérisé en ce que l'on cultive une cellule transformée par un vecteur selon la revendication 59 dans des conditions permettant l'expression dudit polypeptide et que l'on recupère ledit polypeptide recombinant.
- 15 65. Polypeptide recombinant susceptible d'être obtenu par un procédé selon la revendication 64.
  - 66. Procédé de préparation d'un polypeptide synthétique selon l'une des revendications 5 à 7, 29 à 47, caractérisé en ce que l'on effectue une synthèse chimique dudit polypeptide.
- 67. Polypeptide hybride, caractérisé en ce qu'il comprend au moins la séquence d'un polypeptide selon l'une des revendications 5 à 7, 29 à 47 et 65, et une séquence d'un polypeptide susceptible d'induire une réponse immunitaire chez l'homme ou l'animal.
- 68. Séquence nucléotidique codant pour un polypeptide hybride selon la revendication 67.
  - 69. Vecteur caractérisé en ce qu'il contient une séquence nucléotidique selon la revendication 68.
  - 70. Anticorps monoclonal ou polyclonal, ses fragments, ou anticorps chimérique, caractérisé en ce qu'il est capable de reconnaître spécifiquement un polypeptide selon l'une des revendications 5 à 7, 29 à 47, 65 et 67.
  - 71. Anticorps selon la revendication 70, caractérisé en ce qu'il s'agit d'un anticorps marqué.

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- 72 Procédé pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé dans un échantillon biologique, caractérisé en ce qu'il comprend les étapes suivantes :
- a) mise en contact de l'échantillon biologique avec un anticorps selon l'une des revendications 70 et 71;
- b) mise en évidence du complexe antigène-anticorps éventuellement formé.
- 73. Procédé pour la détection de l'expression d'un gène de Streptococcus agalactiae caractérisé en ce que l'on met en contact une souche de Streptococcus agalactiae, avec un anticorps selon la revendication 70 ou 71 et que l'on détecte le complexe antigène/anticorps éventuellement formé.
- 74. Kit ou nécessaire pour la mise en œuvre d'un procédé selon la revendication72 ou 73, caractérisé en ce qu'il comprend les éléments suivants :
- a) un anticorps selon l'une des revendications 70 et 71;
- b) éventuellement, les réactifs pour la constitution du milieu propice à la réaction immunologique;
  - c) éventuellement, les réactifs permettant la mise en évidence des complexes antigèneanticorps produits par la réaction immunologique.
  - 75. Polypeptide selon l'une des revendications 5 à 7, 32 à 47, 65 et 67, ou anticorps selon l'une des revendications 64 et 65, caractérisé en ce qu'il est immobilisé sur un support, notamment une puce à protéine.
  - 76. Puce à protéine, caractérisée en ce qu'elle contient au moins un polypeptide sclon l'une des revendications 5 à 7, 32 à 47, 65 et 67, ou au moins un anticorps selon l'une des revendications 70 et 71, immobilisé sur le support de ladite puce.
  - 77. Puce à protéine selon la revendication 76, caractérisée en ce qu'elle contient en outre au moins un polypeptide de micro-organisme autre que *Streptococcus* agalactiae ou au moins un anticorps dirigé contre un composé de micro-organisme autre que *Streptococcus agalactiae*, immobilisé sur le support de ladite puce.
  - 78. Kit ou nécessaire pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé, caractérisé en ce qu'il comprend une puce à protéine selon l'une des revendications 76 et 77.
  - 79. Kit ou nécessaire pour la détection et/ou l'identification d'un microorganisme, caractérisé en ce qu'il comprend une puce à protéine selon la revendication 77.



- 80. Procédé de détection et/ou d'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé dans un échantillon biologique, caractérisé en ce qu'il met en œuvre une séquence nucléotidique selon l'une des revendications 2 à 4, 8, 9, 11 à 13, 17 à 25, 48 à 52 et 68.
- 5 81. Procédé selon la revendication 80, caractérisé en ce qu'il comporte les étapes suivantes :
  - a) éventuellement, isolement de l'ADN à partir de l'échantillon biologique à analyser, ou obtention d'un ADNc à partir de l'ARN de l'échantillon biologique;
- b) amplification spécifique de l'ADN de bactéries appartenant à l'espèce Streptococcus
   10 agalactiae ou à un micro-organisme associé à l'aide d'au moins une amorce selon l'une des revendications 48 à 52;
  - c) mise en évidence des produits d'amplification.
  - 82. Procédé selon la revendication 80, caractérisé en ce qu'il comprend les étapes suivantes :
- a) mise en contact d'une sonde nucléotidique selon l'une des revendications 48 à 52, avec un échantillon biologique, l'acide nucléique contenu dans l'échantillon biologique ayant, le cas échéant, préalablement été rendu accessible à l'hybridation, dans des conditions permettant l'hybridation de la sonde à l'acide nucléique d'une bactérie appartenant à l'espèce Streptococcus agalactiae ou à un micro-organisme associé;
  - b) mise en évidence de l'hybride éventuellement formé entre la sonde nucléotidique et l'acide nucléique de l'échantillon biologique.
  - 83. Procédé selon la revendication 80, caractérisé en ce qu'il comprend les étapes suivantes :
- a) mise en contact d'une sonde nucléotidique immobilisée sur un support selon la revendication 50 avec un échantillon biologique, l'acide nucléique de l'échantillon ayant, le cas échéant, été préalablement rendu accessible à l'hybridation, dans des conditions permettant l'hybridation de la sonde à l'acide nucléique d'une bactérie appartenant à l'espèce Streptococcus agalactiae ou à un micro-organisme associé;
- b) mise en contact de l'hybride formé entre la sonde nucléotidique immobilisée sur un support et l'acide nucléique contenu dans l'échantillon biologique, le cas échéant après élimination de l'acide nucléique de l'échantillon biologique n'ayant pas hybridé avec la sonde, avec une sonde nucléotidique marquée selon la revendication 49;

- c) mise en évidence du nouvel hybride formé à l'étape b).
- 84. Procédé selon la revendication 83, caractérisé en ce que, préalablement à l'étape a), l'ADN de l'échantillon biologique ou l'ADNc obtenu éventuellement par transcription inverse de l'ARN de l'échantillon, est amplifié à l'aide d'au moins une amorce selon l'une des revendications 48 à 52.
- 85. Kit ou nécessaire pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :
- a) une sonde nucléotidique selon l'une des revendications 48 à 52;
- b) éventuellement, les réactifs nécessaires à la mise en œuvre d'une réaction d'hybridation;
  - c) éventuellement, au moins une amorce selon l'une des revendications 48 à 52 ainsi que les réactifs nécessaires à une réaction d'amplification de l'ADN.
- 86. Kit ou nécessaire pour la détection et/ou l'identification de bactéries appartenant à l'espèce Streptococcus agalactiae ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :
  - a) une sonde nucléotidique, dite sonde de capture, selon la revendication 50;
  - b) une sonde oligonucléotidique, dite sonde de révélation, selon la revendication 49;
- c) éventuellement, au moins une amorce selon l'une des revendications 48 à 52 ainsi que les réactifs nécessaires à une réaction d'amplification de l'ADN.
  - 87. Kit ou nécessaire pour la détection et/ou l'identification de bactéries appartenant à l'espèce *Streptococcus agalactiae* ou à un micro-organisme associé, caractérisé en ce qu'il comprend les éléments suivants :
  - a) au moins une amorce selon l'une des revendications 48 à 52;
- b) éventuellement, les réactifs nécessaires pour effectuer une réaction d'amplification d'ADN;
  - c) éventuellement, un composant permettant de vérifier la séquence du fragment amplifié, plus particulièrement une sonde oligonucléotidique selon l'une des revendications 48 à 52.
- 88. Procédé selon les revendications 72, 73 et 80 à 84 ou kit ou nécessaire selon les revendications 74, 78, 79 et 85 à 87 pour la détection et/ou l'identification de bactéries appartenant à l'espèce Streptococcus agalactiae, caractérisé en ce que ladite amorce et/ou ladite sonde sont choisies parmi les séquences nucléotidiques selon l'une des revendications 2 à 4, 8 à 28, 48 à 52, et 68 spécifiques de l'espèce Streptococcus

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agalactiae, en ce que lesdits polypeptides sont choisis parmi les polypeptides selon l'une des revendications 5 à 7, 32 à 47, 65 et 67 spécifiques de l'espèce Streptococcus agalactiae et en ce que lesdits anticorps sont choisis parmi les anticorps selon l'une des revendications 70 et 71 dirigés contre les polypeptides choisis parmi les polypeptides selon l'une des revendications 5 à 7, 32 à 47, 65 et 67 spécifiques de l'espèce Streptococcus agalactiae.

- 89. Souche de *Streptococcus agalactiae*, caractérisée en ce qu'elle contient au moins une mutation dans au moins une séquence nucléotidique selon l'une des revendications 2 à 4, 8 à 28.
- 90. Souche de *Streptococcus agalactiae* selon la revendication 89, caractérisée en ce que la mutation mène à une inactivation du gène.
  - 91. Souche de *Streptococcus agalactiae* selon la revendication 89, caractérisée en ce que la mutation mène à une surexpression du gène.
  - 92. Utilisation d'une séquence nucléotidique selon l'une des revendications 2 à 4, 8 à 28, d'un polypeptide selon l'une des revendications 5 à 7, 32 à 47, 65 et 67 d'un anticorps selon l'une des revendications 70 et 71, d'une cellule selon l'une des revendications 60 à 62, et/ou d'un animal transformé selon la revendication 63 pour la sélection de composé organique ou inorganique capable de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, et/ou de modifier la réplication cellulaire de cellules eucaryotes ou procaryotes ou capables d'induire, d'inhiber ou d'aggraver chez un organisme animal ou humain les pathologies liées à une infection par Streptococcus agalactiae ou par un micro-organisme associé.
  - 93. Méthode de sélection de composé capable de se lier à un polypeptide selon l'une des revendications 5 à 7, 32 à 47, 65 et 67, capable de se lier à une séquence nucléotidique selon l'une des revendications 2 à 4, 8 à 28, ou capable de reconnaître un anticorps selon l'une des revendications 70 et 71, et/ou capable de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, et/ou de modifier la réplication cellulaire de cellules eucaryotes ou procaryotes, ou capable d'induire, d'inhiber ou d'aggraver chez un organisme animal ou humain les pathologies liées à une infection par Streptococcus agalactiae, caractérisée en ce qu'elle comprend les étapes suivantes :
  - a) mise en contact dudit composé avec ledit polypeptide, ladite séquence nucléotidique, avec une cellule transformée selon l'une des revendications 60 à 62, et/ou administration dudit composé à un animal transformé selon la revendication 63;



- b) détermination de la capacité dudit composé à se lier avec ledit polypeptide ou ladite séquence nucléotidique, ou de moduler, de réguler, d'induire ou d'inhiber l'expression de gènes, ou de moduler la croissance ou la réplication cellulaire, ou d'induire, d'inhiber ou d'aggraver chez ledit organisme animal ou humain les pathologies liées à une infection par Streptococcus agalactiae ou par un microorganisme associé.
- 94. Composition pharmaceutique comprenant un composé choisi parmi les composés suivants :
- a) une séquence nucléotidique selon l'une des revendications 2 à 4, 8 à 28;
- b) un polypeptide selon l'une des revendications 5 à 7, 32 à 47, 65 et 67;
  - c) un vecteur selon la revendication 59 ou 69; et
  - d) un anticorps selon la revendication 70 ou 71.
  - 95. Composition selon la revendication 94, éventuellement en association avec un véhicule pharmaceutiquement acceptable.
- 96. Composition pharmaceutique selon l'une des revendications 94 et 95 pour la prévention et le traitement d'une infection par une bactérie appartenant à l'espèce Streptococcus agalactiae.
  - 97. Composition immunogène, caractérisée en ce qu'elle comprend un ou plusieurs polypeptides selon l'une des revendications 5 à 7, 32 à 47, 65, et/ou un ou plusieurs polypeptides hybrides selon la revendication 67.
  - 98. Utilisation d'une cellule selon l'une des revendications 60 à 62, ou d'un vecteur selon l'une des revendications 59 ou 69 pour la préparation d'une composition vaccinale.
- 99. Composition vaccinale, caractérisée en ce qu'elle contient un polynucléotide selon l'une des revendications 1 à 4, 8 à 28, un vecteur selon l'une des revendications 59 ou 69, et/ou une cellule selon l'une des revendications 60 à 62.
  - 100. Composition vaccinale, caractérisée en ce qu'elle contient au moins un polypeptide codé par un polypucléotide de séquence choisie parmi SEQ ID N°1503,678,2192,1861,584,280.
- 30 101. Composition vaccinale selon la revendication 100, caractérisée en ce qu'il s'agit d'une composition vétérinaire
  - 102. Composition immunogène capable d'induire une réponse immunitaire cellulaire ou humorale pour la prévention ou le traitement d'une infection par une bactérie appartenant à l'espèce Streptococcus agalactiae, caractérisée en ce qu'elle

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comprend une composition immunogène selon la revendication 97, ou une composition vaccinale selon la revendication 99 ou 100, en association avec un véhicule pharmaceutiquement acceptable et éventuellement un ou plusieurs adjuvants de l'immunité appropriés.

- 103. Banque génomique de *Streptococcus agalactiae* CIP 82.45 (ATCC 12403).
- 104. Banque d'ADN génomique selon la revendication 101, caractérisée en ce que ladite banque d'ADN est clonée dans un plasmide.
- 105. Banque selon la revendication 101 ou 102, caractérisée en ce qu'il s'agit de la banque déposée à la CNCM le 28 décembre 2000 sous le N° I-2610.
  - 106. Utilisation des banques génomiques selon l'une des revendications 101 à 103 pour isoler des séquences nucléotidiques spécifiques de *Streptococcus agalactiae*, caractérisée en ce que les séquences nucléotidiques de *Streptococcus* autres que *Streptococcus agalactiae* CIP 82.45 (ATCC 12403) sont alignées et en ce que les données obtenues par cet alignement sont traitées pour isoler lesdites séquences spécifiques.
  - 107. Procédé d'identification de séquence spécifique de Streptococcus agalactiae, caractérisé par l'alignement de séquences nucléotidiques de Streptococcus agalactiae selon les revendications 1 à 4, 8 à 9 et le traitement des données obtenues par cet alignement pour isoler les séquences spécifiques.
- 108. Souche mutante NEM 1979 de *Streptococcus agalactiae* selon la revendication 89 déposée à la CNCM le 24 avril 2002 sous le N° I-2861.
- 109. Souche mutante NEM 2056 de *Streptococcus agalactiae* selon la revendication 89 déposée à la CNCM le 24 avril 2002 sous le N° I-2862.
- 25 110. Souche mutante NEM 2057 de *Streptococcus agalactiae* selon la revendication 89 déposée à la CNCM le 24 avril 2002 sous le N° I-2863.

### (19) Organisation Mondiale de la Propriété Intellectuelle

Bureau international



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(43) Date de la publication internationale 21 novembre 2002 (21.11.2002)

### (10) Numéro de publication internationale WO 02/092818 A3

(51) Classification internationale des brevets7: C12N 15/31, C07K

14/315, 16/12, C12N 15/63, A61K 39/09, 39/40, 31/711, C12Q 1/68, G01N 33/53, A01K 671/027, A01H 5/00

(21) Numéro de la demande internationale :

PCT/IB02/03059

- (22) Date de dépôt international : 26 avril 2002 (26.04.2002)
- (25) Langue de dépôt :

français

(26) Langue de publication :

français

(30) Données relatives à la priorité : 2001/05642 26 avril 2001 (26.04.2001)

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- (81) États désignés (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZM, ZW.
- (84) États désignés (régional): brevet ARIPO (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), brevet eurasien (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), brevet européen (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), brevet OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

#### Déclaration en vertu de la règle 4.17 :

relative à la qualité d'inventeur (règle 4.17.iv)) pour US seulement

### Publiée:

- avec rapport de recherche internationale
- avec la partie réservée au listage des séquences de la description publiée séparément sous forme électronique et disponible sur demande auprès du Bureau international
- (88) Date de publication du rapport de recherche internationale: 28 août 2003

[Suite sur la page suivante]

- (54) Title: STREPTOCOCCUS AGALACTIAE GENOME SEQUENCE, USE FOR DEVELOPING VACCINES, DIAGNOSTIC TOOLS, AND FOR IDENTIFYING THERAPEUTIC TARGETS
- (54) Titre: SEQUENCE DU GENOME STREPTOCOCCUS AGALACTIAE ET SES UTILISATIONS
- (57) Abstract: The invention concerns the genome sequence and nucleotide sequences coding for Streptococcus agalactiae polypeptides, such as cellular envelope polypeptides, or secreted or specific polypeptides, or polypeptides involved in the metabolism and the replication process, as well as vectors or cells comprising said sequences. The invention also concerns the use thereof for developing vaccines, diagnostic tools, DNA chips and for identifying therapeutic targets.
- (57) Abrégé: L'invention a pour objet la séquence génomique et des séquences nucléotidiques codant pour des polypeptides de Streptococcus agalactiae, tels que des polypeptides d'enveloppe cellulaire, ou des polypeptides sécrétés ou spécifiques, ou impliqués dans le métabolisme et dans le processus de réplication, ainsi que des vecteurs ou cellules incluant lesdites séquences. L'invention porte également sur leur application au développement de vaccins, d'outils de diagnostic, de puces à ADN et à l'identification de cibles thérapeutiques.







En ce qui concerne les codes à deux lettres et autres abréviations, se référer aux "Notes explicatives relatives aux codes et abréviations" figurant au début de chaque numéro ordinaire de la Gazette du PCT.

## INTERNATIONAL STARCH REPORT

International approximation No. PCT/IB 02/03059

A. CLASSIFICATION OF SUBJECT MATTER C12N15/31 C07K14/315 C07K16/12 C12N15/63 IPC 7 A61K39/09 A61K39/40 A61K31/711 C12Q1/68 G01N33/53 A01K67/027 According to International Patent Classification (IPC) or to both national classification and IPC A01H5/00

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 C12N C07K A61K

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, WPI Data, PAJ, BIOSIS, CHEM ABS Data, MEDLINE, EMBL

### C. DOCUMENTS CONSIDERED TO BE RELEVANT --

Category •	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
<b>X</b>	SPELLERBERG B ET AL: "LMB, A PROTEIN WITH SIMILARITIES TO THE LRAI ADHESIN FAMILY, MEDIATES ATTACHMENT OF STREPTOCOCCUS AGALACTIAE TO HUMAN LAMININ" INFECTION AND IMMUNITY, AMERICAN SOCIETY FOR MICROBIOLOGY. WASHINGTON, US, vol. 67, no. 2, February 1999 (11.02.99), pages 871-878, XP000973065 ISSN: 0019-9567 -& DATABASE EMBL [en ligne] 11 February 1999 (11.02.99) SPELLERBERG B. ET AL.: "Streptococcus agalactiae Lmb (1mb) gene, complete cds; and unknown gene" Database accession no. AF062533 XP002221154 The whole document	1-9,12, 31, 48-99, 102, 107-110

Ì					
	X	Further documents are listed in the continuation of Box C.	See patent family annex.		
	* "A" "E"	Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance	the principle of theory underlying the invention		
	"L"	earlier document but published on or after the international filing dat document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other	is considered novel or cannot be considered to involve an inventive		
	"O"	means	combined with one or more other such documents, such combination		
		the priority date claimed	"&" document member of the same patent family		
Ì	Date of the actual completion of the international search		Date of mailing of the international search report		
	18 November 2002 (18.11.02)		19 February 2003 (19.02.03)		
	Name and mailing address of the ISA/		Authorized officer		
	S.P.T.O.				
-	Facsimile No.		Telephone No.		

### INTERNATIONAL SEARCH REPORT



ational application No.

C (Continuat	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
X	WO 01 14421 A (MEDIMMUNE, INC.) 1 March 2001 (01.03.01)  page 11, line 13 -page 26, line 20 Sequence listing SEQ ID NO:5, 6	1-9,12, 31, 48-99, 102, 107-110	
X .	DATABASE SWALL [in line]  1 March 2001 (01.03.01)  DE BOEVER, E.H. ET AL.: "TraG-related protein"  Database accession no. Q9F1G0  XP002221155  The whole document  & ERIKA H. DE BOEVER ET AL.: "Enterococcus faecalis conjugative plasmid pAM373: complete nucleotide sequence and genetic analysis of sex phromone response".  MOLECULAR MICROBIOLOGY,  vol. 37, no. 6, 2000, pages 1327-1341,	5,7,8	
A	DORAN, T. I. ET AL.: "Factors Influencing Release of Type III Antigens by Group B Streptococci" INFECT. IMMUN., vol. 31, no. 2, February 1981 (02,02,81) pages 615-623, XP002191322 page 621, hand left column, alinéa 2 - alinéa 3	1-102, 107-110	
		7	

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

## INTERNATIONAL RCH REPORT

į	International appoint No.
	PCT/IB 02/03059

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)			
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:				
1.	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:			
2.	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:			
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).			
Вох П	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)			
This Inte	emational Searching Authority found multiple inventions in this international application, as follows:			
	see supplementary sheet			
1 2	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.  As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.			
3.	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:			
4. X	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:			
	In particular 1-102, 107-110			
Remark	on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.			

Form PCT/ISA/210 (continuation of first sheet (1)) (July 1992)

The International Searching Authority has determined that the present international application contains multiple (groups of) inventions, namely:

1. Claims: (in part) 1-102, 107-110

Nucleotide sequence of Streptococcus agalactiae, SEQ ID No. 1, fragments and homologues; polypeptides coded by said sequences, derived polypeptides, hybrids and antibodies; probe and primer derived from the nucleotide sequence; use thereof for selecting compounds having an effect on disease states caused by an S. agalactiae infection; pharmaceutical compositions; DNA chips and protein; kit containing said chips; cloning vector, host cell, plant or animal containing said nucleotide sequence; use of the antibodies and of the nucleotide sequences to identify Streptococcus agalactiae; strain of S. agalactiae containing a mutation in the sequence SEQ ID No. 1; use of the nucleotide sequences to identify S. agalactiae-specific sequences.

2. Claims: (in part) 1-102, 107-110

Same as invention n° 1, for the nucleotide sequences SEQ ID Nos. 2-139, 2345 and 4482-6617 respectively.

3. Claims: 103-106

Gene library of Streptococcus agalactiae and use thereof.

## INTERNATIONAL ARCH REPORT Information on part mily members

International Appl No
PCT/IB 02/03059

Patent document cited in search report		Publication date	Patent familiy member(s)		Publication date	
WO 0114421	Α.	01-03-2001	AU	7076100 A	19-03-2001	
•			EP	1210366 A1	05-06-2002	
			MO	0114421 A1	01-03-2001	

Form PCT/ISA/210 (patent family annex) (July 1992)

A. CLASSEMENT DE L'OBJET DE LA DEMANDE CIB 7 C12N15/31 C07K14/315

A61K39/40

A01H5/00

A61K31/711

C07K16/12 C12Q1/68

C12N15/63 G01N33/53

A61K39/09 A01K67/027

Selon la classification internationale des brevets (CIB) ou à la fois selon la classification nationale et la CIB

### B. DOMAINES SUR LESQUELS LA RECHERCHE A PORTE

Documentation minimate consultée (système de classification suivi des symboles de classement) C1B 7 C12N C07K A61K

Documentation consultée autre que la documentation minimale dans la mesure où ces documents relèvent des domaines sur lesquels a porté la recherche

Base de données électronique consultée au cours de la recherche internationale (nom de la báse de données, et si réalisable, termes de recherche utilisés) EPO-Internal, WPI Data, PAJ, BIOSIS, CHEM ABS Data, MEDLINE, EMBL

Catégorie °	Identification des documents cités, avec, le cas échéant, l'indication des	des nassanes nertinonts	no. des revendications visées
Categorie	identification des documents dies, avec, le das editeant numeron d	ues passages permients .	110. des revendications visees
X	SPELLERBERG B ET AL: "LMB, A PROSIMILARITIES TO THE LRAI ADHESIN MEDIATES ATTACHMENT OF STREPTOCOCKAGALACTIAE TO HUMAN LAMININ" INFECTION AND IMMUNITY, AMERICAN FOR MICROBIOLOGY. WASHINGTON, US, vol. 67, no. 2, février 1999 (1999 pages 871-878, XP000973065 ISSN: 0019-9567 -& DATABASE EMBL [en ligne] 11 février 1999 (1999-02-11) SPELLERBERG B. ET AL.: "Streptocotagalactiae Lmb (1mb) gene, completand unknown gene" Database accession no. AF062533 XP002221154 le document en entier	FAMILY, CUS SOCIETY 9-02),	1-9,12, 31, 48-99, 102, 107-110
X Voir la	a suite du cadre C pour la fin de la liste des documents	X Les documents de familles de br	evets sont indiqués en annexe
*Catégories spéciales de documents cités:  "A" document définissant l'état général de la technique, non considéré comme particulièrement pertinent  "E" document antérieur, mais publié à la date de dépôt international ou après cette date  "L" document pouvant jeter un doute sur une revendication de priorité ou cité pour déterminer la date de publication d'une autre citation ou pour une raison spéciale (telle qu'indiquée)  "O" document se référant à une divulgation orale, à un usage, à une exposition ou tous autres moyens  "P" document publié avant la date de dépôt international, mais postérieurement à la date de priorité revendiquée  Date à laquelle la recherche internationale a été effectivement achevée  "T" document ultérieur publié après la date de dépôt international date de priorité et n'appartemenant pas à l'état de la technique pertinent, mais cité pour comprendre le principe ou la théorie constituant la base de l'invention revendiquée s'tre considérée comme nouvelle ou comme impliquant une a inventive par rapport au document considéré solément "Y" document particulièrement pertinent; l'inven tion revendiquée s'tre considérée comme nouvelle ou comme impliquant une a l'une autre citation ou pour une activité inventions que de depôt international document particulièrement pertinent; l'inven tion revendiquée s'tre considérée comme nouvelle ou comme impliquant une a divide de priorité de l'appartemenant pas à l'état de la dete de priorité et n'appartemenant pas à l'état de la technique pertinent, mais cité pour che principe ou la théorie constituant la base de l'invention ou l		as à l'état de la ommrendre le principe invention revendiquée ne peut comme impliquant une activité onsidéré isolément l'invention revendiquée iquant une activité inventive nou plusieurs autres ombinaison étant évidente amille de brevets	
		Date d'expédition du présent rapport	
lom et adres	se postale de l'administration chargée de la recherche internationale Office Européen des Brevets, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Fonctionnaire autorisé  MONTERO LOPEZ B.	

## RAPPORT DE RECHERÇ

PCT/IB 02/03059

		PC1/18 02	2/00003
	OCUMENTS CONSIDERES COMME PERTINENTS		
Catégorie <sup>e</sup>	Identification des documents cités, avec, le cas échéant, l'indication des passages p	ertinents	no. des revendications visées
X	WO 01 14421 A (MEDIMMUNE, INC.) 1 mars 2001 (2001-03-01)		1-9,12, 31, 48-99, 102, 107-110
	page 11, ligne 13 -page 26, ligne 20 Sequence listing SEQ ID NO:5, 6		
(	DATABASE SWALL [en ligne] 1 mars 2001 (2001-03-01) DE BOEVER, E.H. ET AL.: "TraG-related protein" Database accession no. Q9F1G0 XP002221155		5,7,8
	le document en entier & ERIKA H. DE BOEVER ET AL.: "Enterococcus faecalis conjugative plasmid pAM373: complete nucleotide sequence and genetic analysis of sex phromone response" MOLECULAR MICROBIOLOGY, vol. 37, no. 6, 2000, pages 1327-1341,		
	DORAN, T. I. ET AL.: "Factors Influencing Release of Type III Antigens by Group B Streptococci" INFECT. IMMUN., vol. 31, no. 2, février 1981 (1981-02), pages 615-623, XP002191322 page 621, colonne de gauche, alinéa 2 - alinéa 3		1-102, 107-110
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Demande internationale n° PCT/IB. 02/03059

Cadre I Observations - lorsqu'îl a été estimé que certaines revendications ne pouvaient pas faire l'objet d'une recherche (suite du point 1 de la première feuille)
Conformément à l'article 17.2)a), certaines revendications n'ont pas fait l'objet d'une recherche pour les motifs suivants:
1. Les revendications n°s se rapportent à un objet à l'égard duquel l'administration n'est pas tenue de procéder à la recherche, à savoir:
2. Les revendications n <sup>os</sup> se rapportent à des parties de la demande internationale qui ne remplissent pas suffisamment les conditions prescrites pour qu'une recherche significative puisse être effectuée, en particulier:
3. Les revendications nos sont des revendications dépendantes et ne sont pas rédigées conformément aux dispositions de la deuxième et de la troisième phrases de la règle 6.4.a).
Cadre II Observations - lorsqu'il y a absence d'unité de l'invention (suite du point 2 de la première feuille)
L'administration chargée de la recherche internationale a trouvé plusieurs inventions dans la demande internationale, à savoir:
voir feuille supplémentaire
Comme toutes les taxes additionnelles ont été payées dans les délais par le déposant, le présent rapport de recherche internationale porte sur toutes les revendications pouvant faire l'objet d'une recherche.
Comme toutes les recherches portant sur les revendications qui s'y prêtalent ont pu être effectuées sans effort particulier justifiant une taxe additionnelle, l'administration n'a sollicité le paiement d'aucune taxe de cette nature.
3. Comme une partie seulement des taxes additionnelles demandées a été payée dans les délais par le déposant, le présent rapport de recherche internationale ne porte que sur les revendications pour lesquelles les taxes ont été payées, à savoir les revendications n os
Aucune taxe additionnelle demandée n'a été payée dans les délais par le déposant. En conséquence, le présent rapport de recherche internationale ne porte que sur l'invention mentionnée en premier lieu dans les revendications; elle est couverte par les revendications n os Partiellement 1-102, 107-110
Remarque quant à la réserve   Les taxes additionnelles étaient accompagnées d'une réserve de la part du déposant  Le paiement des taxes additionnelles n'était assorti d'aucune réserve.

Formulaire PCT/ISA/210 (suite de la première feuille (1)) (Juillet 1998)

### SUITE DES RENSEIGNEMENTS INDIQUES SUR PCT/ISA/ 210

L'administration chargée de la recherche internationale a trouvé plusieurs (groupes d') inventions dans la demande internationale, à savoir:

1. revendications: Partiellement 1-102, 107-110

Séquence nucléotidique de Streptococcus agalactiae SEQ ID NO:1, fragments et homologues; polypeptides codés par ces séquences, polypeptides derivés, hybrides et anticorps; sonde et amorce dérivés de la séquence nucléotidique; utilisation de ceux-ci pour la sélection de composés ayant une influence sur les pathologies liées à une infection par S. agalactiae; compositions pharmaceutiques; puces à ADN et protéine; kit ou nécessaire contenant lesdites puces; vecteur de clonage, cellule hôte, végétal ou animal contenant la séquence nucléotidique; utilisation des anticorps et des séquences nucléotidiques pour l'identification de Streptococcus agalactiae; souche de S. agalactiae contenant une mutation dans la séquence SEQ ID NO:1; utilisation des séquences nucléotidiques pour identifier des séquences spécifiques de S. agalactiae.

2. revendications: Partiellement 1-102, 107-110

Idem au sujet 1 pour, respectivement les séquences nucléotidiques SEQ ID NOs:2-139, 2345, and 4482-6617

3. revendications: 103-106

Banque génomique de Straptococcus agalactiae et son utilisation

### PPORT DE RECHERCHE INTERNATIONALE

mationale No bres de familles de brevets Renseignements relatifs a 02/03059 Document brevet cité au rapport de recherche Date de publication Membre(s) de la famille de brevet(s) Date de publication 19-03-2001 Α 01-03-2001 ΑU 7076100 A WO 0114421 05-06-2002 01-03-2001 ΕP 1210366 A1 WO 0114421 A1

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